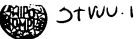
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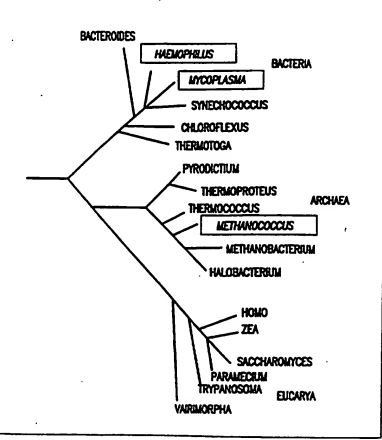
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(57) Abstract

The present application describes the complete 1.66-megabase pair genome sequence of an autotrophic archaeon, *Methanococcus jannaschii*, and its 58- and 16-kilobase pair extrachromosomal elements. Also described are 1738 predicted proteincoding genes.



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C mplete Genome Sequence of the Methan genic Archaeon, Methanococcus jannaschii

Background of the Invention

Statement as to Rights to Inventions Made Under Federally-Sponsored Research and Development

Part of the work performed during development of this invention utilized U.S. Government funds. The U.S. Government may have certain rights in the invention - DE-FC02-95ER61962; DE-FC02-95ER61963; and NAGW 2554.

Field of the Invention

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The present application discloses the complete 1.66-megabase pair genome sequence of an autotrophic archaeon, *Methanococcus jannaschii*, and its 58- and 16-kilobase pair extrachromosomal elements. Also identified are 1738 predicted protein-coding genes.

Related Background Art

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The view of evolution in which all cellular organisms are in the first instance either prokaryotic or eukaryotic was challenged in 1977 by the finding that on the molecular level life comprises three primary groupings (Fox, G.E., et al., Proc. Natl. Acad. Sci. USA 74:4537 (1977); Woese, C.R. & Fox, G.E., Proc. Natl. Acad. Sci. USA 74:5088 (1977); Woese, C.R., et al., Proc. Natl. Acad. Sci. USA 87:4576 (1990)): the eukaryotes (Eukarya) and two unrelated groups of prokaryotes, Bacteria and a new group now called the Archaea. Although Bacteria and Archaea are both prokaryotes in a cytological sense, they differ profoundly in their molecular makeup (Fox, G.E., et al., Proc. Natl. Acad. Sci. USA 74:4537 (1977); Woese, C.R. & Fox, G.E., Proc. Natl. Acad. Sci. USA 74:5088 (1977); Woese, C.R., et al., Proc. Natl. Acad. Sci. USA 74:5088 (1977); Woese, C.R., et al., Proc. Natl. Acad. Sci. USA 74:5088 (1977); Woese, C.R., et al., Proc. Natl. Acad. Sci. USA 74:5088 (1977); Woese, C.R., et al., Proc. Natl. Acad. Sci. USA 87:4576 (1990)).

Several lines of molecular evidence even suggest a specific relationship between Archaea and Eukarya (Iwabe, N., et al., Proc. Natl. Acad. Sci. USA 86:9355 (1989); Gogarten J.P., et al., Proc. Natl. Acad. Sci. USA 86:6661 (1989); Brown, J.R. and Doolittle, W.F., Proc. Natl. Acad. Sci. USA 92:2441 (1995)).

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The era of true comparative genomics has been ushered in by complete genome sequencing and analysis. We recently described the first two complete bacterial genome sequences, those of *Haemophilus influenzae* and *Mycoplasma genitalium* (Fleischmann, R.D., et al., Science 269:496 (1995); Fraser, C.M., et al., Science 270:397 (1995)). Large scale DNA sequencing efforts also have produced an extensive collection of sequence data from eukaryotes, including *Homo sapiens* (Adams, M.D., et al., Nature 377:3 (1995)) and Saccharomyces cerevisiae (Levy, J., Yeast 10:1689 (1994)).

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M. jannaschii was originally isolated by J.A. Leigh from a sediment sample collected from the sea floor surface at the base of a 2600 m deep "white smoker" chimney located at 21°N on the East Pacific Rise (Jones, W., et al., Arch. Microbiol. 136:254 (1983)). M. jannaschii grows at pressures of up to more than 500 atm and over a temperature range of 48-94 °C, with an optimum temperature near 85 °C (Jones, W., et al., Arch. Microbiol. 136:254 (1983)). The organism is autotrophic and a strict anaerobe; and, as the name implies, it produces methane. The dearth of archaeal nucleotide sequence data has hampered attempts to begin constructing a comprehensive comparative evolutionary framework for assessing the molecular basis of the origin and diversification of cellular life.

Summary of the Invention

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The present invention is based on whole-genome random sequencing of an autotrophic archaeon, *Methanococcus jannaschii*. The *M. jannaschii* genome consists of three physically distinct elements: (i) a large circular chromosome; (ii) a large circular extrachromosomal element (ECE); and (iii) a small circular extrachromosomal element (ECE). The nucleotide sequences generated, the *M*.

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jannaschii chromosome, the large ECE, and the small ECE, are respectively provided on pages 152-585 (SEQ ID NO:1), pages 585-600 (SEQ ID NO:2), and pages 601-605 (SEQ ID NO:3).

The present invention is further directed to isolated nucleic acid molecules comprising open reading frames (ORFs) encoding *M. jannaschii* proteins. The present invention also relates to variants of the nucleic acid molecules of the present invention, which encode portions, analogs or derivatives of *M. jannaschii* proteins. Further embodiments include isolated nucleic acid molecules comprising a polynucleotide having a nucleotide sequence at least 90% identical, and more preferably at least 95%, 96%, 97%, 98% or 99% identical, to the nucleotide sequence of a *M. jannaschii* ORF described herein.

The present invention also relates to recombinant vectors, which include the isolated nucleic acid molecules of the present invention, host cells containing the recombinant vectors, as well as methods for making such vectors and host cells for *M. jannaschii* protein production by recombinant techniques.

The invention further provides isolated polypeptides encoded by the *M. jannaschii* ORFs. It will be recognized that some amino acid sequences of the polypeptides described herein can be varied without significant effect on the structure or function of the protein. If such differences in sequence are contemplated, it should be remembered that there will be critical areas on the protein which determine activity. In general, it is possible to replace residues which form the tertiary structure, provided that residues performing a similar function are used. In other instances, the type of residue may be completely unimportant if the alteration occurs at a non-critical region of the protein.

In another aspect, the invention provides a peptide or polypeptide comprising an epitope-bearing portion of a polypeptide of the invention. The epitope-bearing portion is an immunogenic or antigenic epitope useful for raising antibodies.

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Brief Description of the Figures

Figure 1. A schematic showing the relationship of the three domains of life based on sequence data from the small subunit of rRNA (Fox, G.E., et al., Proc. Natl. Acad. Sci. USA 74:4537 (1977); Woese, C.R. & Fox, G.E., Proc. Natl. Acad. Sci. USA 74:5088 (1977); Woese, C.R., et al., Proc. Natl., Acad. Sci. USA 87:4576 (1990)).

Figure 2. Structure of a putative family of insertion sequence (IS) elements in the M. jannaschii genome. The family of elements has been named ISAMJI and contains 11 members distributed among three groups (A, B, and C). The outer rectangle indicates the entire IS element; the interior rectangles indicate the predicted coding regions, oriented with the NH2-termini to the left. DNA immediately adjacent to the NH₂-termini is 75 to 100% identical over 50 bp; DNA sequence similarity at the COOH-termini ends immediately after the stop codon. Black triangles indicate terminal inverted repeats. Fill patterns indicate which regions are missing from the elements in groups B and C. (A) Two copies of this family are 642 bp long and are 97% similar to each other at the nucleotide level. They appear to encode a protein 214 amino acids in length (ORFs MJ0017 and MJ1466) that are 27% identical to the IS240 transposase of Bacillus thuriengiensis (GenBank Accession number: M23741). (B) Eight copies of the family range in length from 358 to 360 bp and are missing a 342-bp internal region relative to the two members of group A. Some members of group B have putative frameshifts (indicated by solid arrows) and in-frame UGA codons (indicated by open arrows). (C) The single copy in group C is 265 bp in length and occurs on the large ECE. The 436 bp internal region missing from this element is different than that of the members of group B.

Figure 3. Structure of a multicopy repetitive element in the *M. jannaschii* genome. Of the 18 copies identified on the main chromosome, seven are oriented in one direction (plus strand) and 11 are oriented in the opposite strand. Each element consists of a long, 391- to 425-bp repeat segment (designated LR) followed by up to 25 short, 27- to 28-bp repeat segments (designated SR). Each

SR segment is separated by 31 to 51 bp of sequence that is unique within and between each complete repeat lement. (A) The longest repeat element has an LR segment followed by 25 SR segments, and spans more than 2 kbp, and (B) the shortest complete element has an LR segment followed by two SR segments. (C) One element is present in the genome with five SR segments and no LR component. (D and E) The LR segments of two elements in the genome are truncated at the end adjacent to the SR segments, both are followed by a single SR segment.

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Figure 4. Block diagram of a computer system 102 that can be used to implement the computer-based systems of present invention.

Detailed Description of the Invention

The present invention is based on whole-genome random sequencing of an autotrophic archaeon, *Methanococcus jannaschii*. The *M. jannaschii* genome consists of three physically distinct elements: (i) a large circular chromosome of 1,664,976 base pairs (bp) (shown on pages 152-585 and in SEQ ID NO:1), which contains 1682 predicted protein-coding regions and has a G+C content of 31.4%; (ii) a large circular extrachromosomal element (the large ECE) of 58,407 bp (shown on pages 585-600 and in SEQ ID NO:2), which contains 44 predicted protein-coding regions and has a G+C content of 28.2%; and (iii) a small circular extrachromosomal element (the small ECE) of 16,550 bp (shown on pages 601-605 and in SEQ ID NO:3), which contains 12 predicted protein-coding regions and has a G+C content of 28.8%.

The primary nucleotide sequences generated, the *M. jannaschii* chromosome, the large ECE, and the small ECE, are provided in SEQ ID NOs:1, 2, and 3, respectively. As used herein, the "primary sequence" refers to the nucleotide sequence represented by the IUPAC nomenclature system. The present invention provides the nucleotide sequences of SEQ ID NOs:1, 2, and 3, or a representative fragment thereof, in a form which can be readily used, analyzed, and interpreted by a skilled artisan.

As used herein, a "representative fragment" refers to *M. jannaschii* protein-encoding regions (also referred to herein as open reading frames), expression modulating fragments, uptake modulating fragments, and fragments that can be used to diagnose the presence of *M. jannaschii* in a sample. A non-limiting identification of such representative fragments is provided in Tables 2(a) and 3. As described in detail below, representative fragments of the present invention further include nucleic acid molecules having a nucleotide sequence at least 90% identical, preferably at least 95, 96%, 97%, 98%, or 99% identical, to an ORF identified in Table 2(a) or 3.

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As indicated above, the nucleotide sequence information provided in SEQ ID NOs:1, 2 and 3 was obtained by sequencing the M. jannaschii genome using a megabase shotgun sequencing method. The sequences provided in SEQ ID NOs:1, 2 and 3 are highly accurate, although not necessarily a 100% perfect, representation of the nucleotide sequence of the M. jannaschii genome. As discussed in detail below, using the information provided in SEO ID NOs:1, 2 and 3 and in Tables 2(a) and 3 together with routine cloning and sequencing methods, one of ordinary skill in the art would be able to clone and sequence all "representative fragments" of interest including open reading frames (ORFs) encoding a large variety of M. jannaschii proteins. In rare instances, this may reveal a nucleotide sequence error present in the nucleotide sequences disclosed in SEQ ID NOs: 1, 2, and 3. Thus, once the present invention is made available (i.e., once the information in SEQ ID NOs:1, 2, and 3 and in Tables 2(a) and 3 have been made available), resolving a rare sequencing error would be well within the skill of the art. Nucleotide sequence editing software is publicly available. For example, Applied Biosystem's (AB) AutoAssembler™ can be used as an aid during visual inspection of nucleotide sequences.

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Even if all of the rare sequencing errors were corrected, it is predicted that the resulting nucleotide sequences would still be at least about 99.9% identical to the reference nucleotide sequences in SEQ ID NOs:1, 2, and 3. Thus, the present invention further provides nucleotide sequences that are at least 99.9% identical to the nucleotide sequence of SEQ ID NO:1, 2, or 3 in a form which can

be readily used, analyzed and interpreted by the skilled artisan. Methods for determining whether a nucleotide sequence is at least 99.9% identical to a reference nucleotide sequence of the present invention are described below.

Nucleic Acid Molecules

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The present invention is directed to isolated nucleic acid fragments of the *M. jannaschii* genome. Such fragments include, but are not limited to, nucleic acid molecules encoding polypeptides (hereinafter open reading frames (ORFs)), nucleic acid molecules that modulate the expression of an operably linked ORF (hereinafter expression modulating fragments (EMFs)), nucleic acid molecules that mediate the uptake of a linked DNA fragment into a cell (hereinafter uptake modulating fragments (UMFs)), and nucleic acid molecules that can be used to diagnose the presence of *M. jannaschii* in a sample (hereinafter diagnostic fragments (DFs)).

By "isolated nucleic acid molecule(s)" is intended a nucleic acid

molecule, DNA or RNA, that has been removed from its native environment. For

example, recombinant DNA molecules contained in a vector are considered isolated for the purposes of the present invention. Further examples of isolated

DNA molecules include recombinant DNA molecules maintained in heterologous host cells, purified (partially or substantially) DNA molecules in solution, and

nucleic acid molecules produced synthetically. Isolated RNA molecules include

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In one embodiment, M. jannaschii DNA can be mechanically sheared to produce fragments about 15-20 kb in length, which can be used to generate a M. jannaschii DNA library by insertion into lambda clones as described in Example 1 below. Primers flanking an ORF described in Table 2(a) or 3 can then be generated using the nucleotide sequence information provided in SEQ ID NO:1, 2, or 3. The polymerase chain reaction (PCR) is then used to amplify and isolate the ORF from the lambda DNA library. PCR cloning is well known in the art. Thus, given SEQ ID NO:1, 2, and 3, and Tables 2(a) and 3, it would be routine

to isolate any ORF or ther representative fragment of the *M. jannaschii* genome. Isolated nucleic acid molecules of the present invention include, but are not limited to, single stranded and double stranded DNA, and single stranded RNA, and complements thereof.

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Tables 2(a), 2(b) and 3 describe ORFs in the M. jannaschii genome. In particular, Table 2(a) (pages 67-115 below) indicates the location of ORFs (i.e., the position) within the M. jannaschii genome that putatively encode the recited protein based on homology matching with protein sequences from the organism appearing in parentheticals (see the fourth column of Table 2(a)). The first column of Table 2(a) provides a name for each ORF. The second and third columns in Table 2(a) indicate an ORF's position in the nucleotide sequence provided in SEQ ID NO:1, 2 or 3. One of ordinary skill in the art will appreciate that the ORFs may be oriented in opposite directions in the M. jannaschii genome. This is reflected in columns 2 and 3. The fifth column of Table 2(a) indicates the percent identity of the protein sequence encoded by an ORF to the corresponding protein sequence from the organism appearing in parentheticals in the fourth column. The sixth column of Table 2(a) indicates the percent similarity of the protein sequence encoded by an ORF to the corresponding protein sequence from the organism appearing in parentheticals in the fourth column. The concepts of percent identity and percent similarity of two polypeptide sequences are well understood in the art and are described in more detail below. The eighth column in Table 2(a) indicates the length of the ORF in nucleotides. Each identified gene has been assigned a putative cellular role category adapted from Riley (Riley, M., Microbiol. Rev. 57:862 (1993)).

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Table 2(b) (page 116 below) provides the single ORF identified by the present inventors that matches a previously published *M. jannaschii* gene. In particular, ORF MJ0479, which is 585 nucleotides in length and is positioned at nucleotides 1,050,508 to 1,049,948 in SEQ ID NO:1, shares 100% identity to the previously published *M. jannaschii* adenylate kinase gene.

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Table 3 (pages 117-150 below) provides ORFs of the *M. jannaschii* genome that did not elicit a homology match with a known sequence from either

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M. jannaschii r another rganism. As above, the first column in Table 3 provides the ORF name and the second and third columns indicate an ORF's position in SEQ ID NO:1, 2, or 3.

Table 4 (page 151 below) provides genes of M. jannaschii that contain inteins.

In the above-described Tables, there are three groups of ORF names. The one thousand six hundred and eighty two ORFs named "MJ-" (MJ0001-MJ1682) were identified on the *M. jannaschii* chromosome (SEQ ID NO:1). The forty four ORFs named "MJECL-" (MJECL01-MJECL44) were identified on the large ECE (SEQ ID NO:2). The twelve ORFs named "MJECS-" (MJECS01-MJES12) were identified on the small ECE (SEQ ID NO:3).

Further details concerning the algorithms and criteria used for homology searches are provided in the Examples below. A skilled artisan can readily identify ORFs in the *Methanococcus jannaschii* genome other than those listed in Tables 2(a), 2(b) and 3, such as ORFs that are overlapping or encoded by the opposite strand of an identified ORF in addition to those ascertainable using the computer-based systems of the present invention.

Isolated nucleic acid molecules of the present invention include DNA molecules having a nucleotide sequence substantially different than the nucleotide sequence of an ORF described in Table 2(a) or 3, but which, due to the degeneracy of the genetic code, still encode a *M. jannaschii* protein. The genetic code is well known in the art. Thus, it would be routine to generate such degenerate variants.

The present invention further relates to variants of the nucleic acid molecules of the present invention, which encode portions, analogs or derivatives of a *M. Jannaschii* protein encoded by an ORF described in Table 2(a) or 3. Non-naturally occurring variants may be produced using art-known mutagenesis techniques and include those produced by nucleotide substitutions, deletions or additions. The substitutions, deletions or additions may involve one or more nucleotides. The variants may be altered in coding regions, non-coding regions, or both. Alterations in the coding regions may produce conservative or

non-conservative amino acid substitutions, deletions or additions. Especially preferred among these are silent substitutions, additions and deletions, which do not alter the properties and activities of the *M. jannaschii* protein or portions thereof. Also especially preferred in this regard are conservative substitutions.

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As a practical matter, whether any particular nucleic acid molecule is at least 90%, 95%, 96%, 97%, 98% or 99% identical to the nucleotide sequence of a *M. jannaschii* ORF can be determined conventionally using known computer programs such as the Bestfit program (Wisconsin Sequence Analysis Package, Version 8 for Unix, Genetics Computer Group, University Research Park, 575 Science Drive, Madison, WI 53711). Bestfit uses the local homology algorithm

Further embodiments of the invention include isolated nucleic acid molecules comprising a polynucleotide having a nucleotide sequence at least 90% identical, and more preferably at least 95%, 96%, 97%, 98% or 99% identical, to (a) the nucleotide sequence of an ORF described in Table 2(a) or 3, (b) the nucleotide sequence of an ORF described in Table 2(a) or 3, but lacking the codon for the N-terminal methionine residue, if present, or (c) a nucleotide sequence complementary to any of the nucleotide sequences in (a) or (b). By a polynucleotide having a nucleotide sequence at least, for example, 95% identical to the reference M. jannaschii ORF nucleotide sequence is intended that the nucleotide sequence of the polynucleotide is identical to the reference sequence except that the polynucleotide sequence may include up to five point mutations per each 100 nucleotides of the ORF sequence. In other words, to obtain a polynucleotide having a nucleotide sequence at least 95% identical to a reference ORF nucleotide sequence, up to 5% of the nucleotides in the reference sequence may be deleted or substituted with another nucleotide, or a number of nucleotides up to 5% of the total nucleotides in the reference sequence may be inserted into the reference sequence. These mutations of the reference sequence may occur at the 5' or 3' terminal positions of the reference nucleotide sequence or anywhere between those terminal positions, interspersed either individually among nucleotides in the reference sequence or in one or more contiguous groups within the reference sequence.

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of Smith and Waterman, Advances in Applied Mathematics 2: 482-489 (1981), to find the best segment of homology between two sequences. When using Bestfit or any other sequence alignment program to determine whether a particular sequence is, for instance, 95% identical to a reference sequence according to the present invention, the parameters are set, of course, such that the percentage of identity is calculated over the full length of the reference nucleotide sequence and that gaps in homology of up to 5% of the total number of nucleotides in the reference sequence are allowed.

Preferred are nucleic acid molecules having sequences at least 90%, 95%, 96%, 97%, 98% or 99% identical to the nucleic acid sequence of a *M. jannaschii* ORF that encode a functional polypeptide. By a "functional polypeptide" is intended a polypeptide exhibiting activity similar, but not necessarily identical, to an activity of the protein encoded by the *M. jannaschii* ORF. For example, the *M. jannaschii* ORF MJ1434 encodes an endonuclease that degrades DNA. Thus, a "functional polypeptide" encoded by a nucleic acid molecule having a nucleotide sequence, for example, 95% identical to the nucleotide sequence of MJ1434, will also degrade DNA. As the skilled artisan will appreciate, assays for determining whether a particular polypeptide is "functional" will depend on which ORF is used as the reference sequence. Depending on the reference ORF, the assay chosen for measuring polypeptide activity will be readily apparent in light of the role categories provided in Table 2(a).

Of course, due to the degeneracy of the genetic code, one of ordinary skill in the art will immediately recognize that a large number of the nucleic acid molecules having a sequence at least 90%, 95%, 96%, 97%, 98%, or 99% identical to the nucleic acid sequence of a reference ORF will encode a functional polypeptide. In fact, since degenerate variants all encode the same amino acid sequence, this will be clear to the skilled artisan even without performing a comparison assay for protein activity. It will be further recognized in the art that, for such nucleic acid molecules that are not degenerate variants, a reasonable number will also encode a functional polypeptide. This is because the skilled artisan is fully aware of amino acid substitutions that are either less likely or not

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likely to significantly affect protein function (e.g., replacing ne aliphatic amino acid with a second aliphatic amino acid).

For example, guidance concerning how to make phenotypically silent amino acid substitutions is provided in Bowie, J. U. et al., "Deciphering the Message in Protein Sequences: Tolerance to Amino Acid Substitutions," Science 247:1306-1310 (1990), wherein the authors indicate that there are two main approaches for studying the tolerance of an amino acid sequence to change. The first method relies on the process of evolution, in which mutations are either accepted or rejected by natural selection. The second approach uses genetic engineering to introduce amino acid changes at specific positions of a cloned gene and selections or screens to identify sequences that maintain functionality. As the authors state, these studies have revealed that proteins are surprisingly tolerant of amino acid substitutions. The authors further indicate which amino acid changes are likely to be permissive at a certain position of the protein. For example, most buried amino acid residues require nonpolar side chains, whereas few features of surface side chains are generally conserved. Other such phenotypically silent substitutions are described in Bowie, J.U. et al., supra, and the references cited therein.

The present invention is further directed to fragments of the isolated nucleic acid molecules described herein. By a fragment of an isolated nucleic acid molecule having the nucleotide sequence of a *M. jannaschii* ORF is intended fragments at least about 15 nt, and more preferably at least about 20 nt, still more preferably at least about 30 nt, and even more preferably, at least about 40 nt in length that are useful as diagnostic probes and primers as discussed herein. Of course, larger fragments 50-500 nt in length are also useful according to the present invention as are fragments corresponding to most, if not all, of the nucleotide sequence of a *M. jannaschii* ORF. By a fragment at least 20 nt in length, for example, is intended fragments that include 20 or more contiguous bases from the nucleotide sequence of a *M. jannaschii* ORF. Since *M. jannaschii* ORFs are listed in Tables 2(a) and 3 and the genome sequence has been provided, generating such DNA fragments would be routine to the skilled artisan. For

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example, restriction endonuclease cleavage or shearing by sonication could easily be used to generate fragments of various sizes. Alternatively, such fragments could be generated synthetically.

Preferred nucleic acid fragments of the present invention include nucleic acid molecules encoding epitope-bearing portions of a *M. jannaschii* protein. Methods for determining such epitope-bearing portions are described in detail below.

In another aspect, the invention provides an isolated nucleic acid molecule comprising a polynucleotide that hybridizes under stringent hybridization conditions to a portion of the polynucleotide in a nucleic acid molecule of the invention described above, for instance, an ORF described in Table 2(a) or 3. By "stringent hybridization conditions" is intended overnight incubation at 42°C in a solution comprising: 50% formamide, 5x SSC (150 mM NaCl, 15mM trisodium citrate), 50 mM sodium phosphate (pH 7.6), 5x Denhardt's solution, 10% dextran sulfate, and 20 g/ml denatured, sheared salmon sperm DNA, followed by washing the filters in 0.1x SSC at about 65°C.

By a polynucleotide that hybridizes to a "portion" of a polynucleotide is intended a polynucleotide (either DNA or RNA) hybridizing to at least about 15 nucleotides (nt), and more preferably at least about 20 nt, still more preferably at least about 30 nt, and even more preferably about 30-70 nt of the reference polynucleotide. These are useful as diagnostic probes and primers as discussed above and in more detail below.

Of course, polynucleotides hybridizing to a larger portion of the reference polynucleotide (e.g., a *M. jannaschii* ORF), for instance, a portion 50-500 nt in length, or even to the entire length of the reference polynucleotide, are also useful as probes according to the present invention, as are polynucleotides corresponding to most, if not all, of a *M. jannaschii* ORF.

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By "expression modulating fragment" (EMF), is intended a series of nucleotides that modulate the expression of an operably linked ORF or EMF. A sequence is said to "modulate the expression of an operably linked sequence" when the expression of the sequence is altered by the presence of the EMF. EMFs include, but are not limited to, promoters, and promoter modulating sequences (inducible elements). One class of EMFs are fragments that induce the expression of an operably linked ORF in response to a specific regulatory factor or physiological event. EMF sequences can be identified within the M. iannaschii genome by their proximity to the ORFs described in Tables 2(a), 2(b), and 3. An intergenic segment, or a fragment of the intergenic segment, from about 10 to 200 nucleotides in length, taken 5' from any one of the ORFs of Tables 2(a), 2(b) or 3 will modulate the expression of an operably linked 3' ORF in a fashion similar to that found with the naturally linked ORF sequence. As used herein, an "intergenic segment" refers to the fragments of the M. jannaschii genome that are between two ORF(s) herein described. Alternatively, EMFs can be identified using known EMFs as a target sequence or target motif in the computer-based systems of the present invention.

The presence and activity of an EMF can be confirmed using an EMF trap vector. An EMF trap vector contains a cloning site 5' to a marker sequence. A marker sequence encodes an identifiable phenotype, such as antibiotic resistance or a complementing nutrition auxotrophic factor, which can be identified or assayed when the EMF trap vector is placed within an appropriate host under appropriate conditions. As described above, an EMF will modulate the expression of an operably linked marker sequence. A more detailed discussion of various marker sequences is provided below.

A sequence that is suspected as being an EMF is cloned in all three reading frames in one or more restriction sites upstream from the marker sequence in the EMF trap vector. The vector is then transformed into an appropriate host using known procedures and the phenotype of the transformed host in examined under appropriate c nditions. As described above, an EMF will modulate the expression of an operably linked marker sequence.

By "uptake modulating fragment" (UMF), is intended a series of nucleotides that mediate the uptake of a linked DNA fragment into a cell. UMFs can be readily identified using known UMFs as a target sequence or target motif with the computer-based systems described below. The presence and activity of a UMF can be confirmed by attaching the suspected UMF to a marker sequence. The resulting nucleic acid molecule is then incubated with an appropriate host under appropriate conditions and the uptake of the marker sequence is determined. As described above, a UMF will increase the frequency of uptake of a linked marker sequence.

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By a "diagnostic fragment" (DF), is intended a series of nucleotides that selectively hybridize to *M. jannaschii* sequences. DFs can be readily identified by identifying unique sequences within the *M. jannaschii* genome, or by generating and testing probes or amplification primers consisting of the DF sequence in an appropriate diagnostic format for amplification or hybridization selectivity.

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Each of the ORFs of the *M. jannaschii* genome disclosed in Tables 2(a) and 3, and the EMF found 5' to the ORF, can be used in numerous ways as polynucleotide reagents. The sequences can be used as diagnostic probes or diagnostic amplification primers to detect the presence *M. jannaschii* in a sample. This is especially the case with the fragments or ORFs of Table 3, which will be highly selective for *M. jannaschii*.

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In addition, the fragments of the present invention, as broadly described, can be used to control gene expression through triple helix formation or antisense DNA or RNA, both of which methods are based on the binding of a polynucleotide sequence to DNA or RNA. Polynucleotides suitable for use in these methods are usually 20 to 40 bases in length and are designed to be complementary to a region of the gene involved in transcription (triple helix - see Lee et al., Nucl. Acids Res. 6:3073 (1979); Cooney et al., Science 241:456 (1988); and Dervan et al., Science 251:1360 (1991)) or to the mRNA itself (antisense - Okano, J. Neurochem. 56:560 (1991); Oligodeoxynucleotides as Antisense Inhibitors of Gene Expression, CRC Press, Boca Raton, FL (1988)).

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Triple helix- formation optimally results in a shut-off of RNA transcription from DNA, while antisense RNA hybridization blocks translation of an mRNA molecule into polypeptide. Both techniques have been demonstrated to be effective in model systems. Information contained in the sequences of the present invention is necessary for the design of an antisense or triple helix oligonucleotide.

Vectors and Host Cells

The present invention further provides recombinant constructs comprising one or more fragments of the *M. jannaschii* genome. The recombinant constructs of the present invention comprise a vector, such as a plasmid or viral vector, into which, for example, a *M. jannaschii* ORF is inserted. The vector may further comprise regulatory sequences, including for example, a promoter, operably linked to the ORF. For vectors comprising the EMFs and UMFs of the present invention, the vector may further comprise a marker sequence or heterologous ORF operably linked to the EMF or UMF. Large numbers of suitable vectors and promoters are known to those of skill in the art and are commercially available for generating the recombinant constructs of the present invention. The following vectors are provided by way of example. Bacterial: pBs, phagescript, PsiX174, pBluescript SK, pBs KS, pNH8a, pNH16a, pNH18a, pNH46a (Stratagene); pTrc99A, pKK223-3, pKK233-3, pDR540, pRIT5 (Pharmacia). Eukaryotic: pWLneo, pSV2cat, pOG44, pXT1, pSG (Stratagene) pSVK3, pBPV, pMSG, pSVL (Pharmacia).

Promoter regions can be selected from any desired gene using CAT (chloramphenicol transferase) vectors or other vectors with selectable markers. Two appropriate vectors are pKK232-8 and pCM7. Particular named bacterial promoters include lacI, lacZ, T3, T7, gpt, lambda P_R, and trc. Eukaryotic promoters include CMV immediate early, HSV thymidine kinase, early and late SV40, LTRs from retrovirus, and mouse metallothionein-I. Selection of the

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appropriate vector and promoter is well within the level fordinary skill in the art.

The present invention further provides host cells containing any one of the isolated fragments (preferably an ORF) of the *M. jannaschii* genome described herein. The host cell can be a higher eukaryotic host cell, such as a mammalian cell, a lower eukaryotic host cell, such as a yeast cell, or the host cell can be a procaryotic cell, such as a bacterial cell. Introduction of the recombinant construct into the host cell can be effected by calcium phosphate transfection, DEAE, dextran mediated transfection, or electroporation (Davis, L. et al., Basic Methods in Molecular Biology (1986)). Host cells containing, for example, a M. jannaschii ORF can be used conventionally to produce the encoded protein.

Polypeptides and Fragments

The invention further provides an isolated polypeptide encoded by a *M. jannaschii* ORF described in Tables 2(a) or 3, or a peptide or polypeptide comprising a portion of the isolated polypeptide. The terms "peptide" and "oligopeptide" are considered synonymous (as is commonly recognized) and each term can be used interchangeably as the context requires to indicate a chain of at least two amino acids coupled by peptidyl linkages. The word "polypeptide" is used herein for chains containing more than ten amino acid residues.

It will be recognized in the art that some amino acid sequence of the *M. jannaschii* polypeptide can be varied without significant affect of the structure or function of the protein. If such differences in sequence are contemplated, it should be remembered that there will be critical areas on the protein which determine activity. In general, it is possible to replace residues which form the tertiary structure, provided that residues performing a similar function are used. In other instances, the type of residue may be completely unimportant if the alteration occurs at a non-critical region of the protein.

Thus, the invention further includes variations of a M. jannaschii protein encoded by an ORF described in Table 2(a) or 3 that show substantial protein

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activity. Methods for assaying such "functional polypeptides" for protein activity are described above. Variations include deletions, insertions, inversions, repeats. and type substitutions (for example, substituting one hydrophilic residue for another, but not strongly hydrophilic for strongly hydrophobic as a rule). Small changes or such "neutral" amino acid substitutions will generally have little effect on protein activity.

Typically seen as conservative substitutions are the replacements, one for another, among the aliphatic amino acids Ala, Val, Leu and Ile; interchange of the hydroxyl residues Ser and Thr, exchange of the acidic residues Asp and Glu, substitution between the amide residues Asn and Gln, exchange of the basic residues Lys and Arg and replacements among the aromatic residues Phe, Tyr.

As indicated in detail above, further guidance concerning amino acid changes that are likely to be phenotypically silent (i.e., are not likely to have a significant deleterious effect on function) can be found in Bowie, J.U., et al., "Deciphering the Message in Protein Sequences: Tolerance to Amino Acid Substitutions," Science 247:1306-1310 (1990).

The fragment, derivative, variant or analog of a *M. jannaschii* polypeptide encoded by an ORF described in Table 2(a) or 3, may be (i) one in which one or more of the amino acid residues are substituted with a conserved or non-conserved amino acid residue (preferably a conserved amino acid residue) and such substituted amino acid residue may or may not be one encoded by the genetic code, or (ii) one in which one or more of the amino acid residues includes a substituent group, or (iii) one in which the polypeptide is fused with another compound, such as a compound to increase the half-life of the polypeptide (for example, polyethylene glycol), or (iv) one in which the additional amino acids are fused to the polypeptide, such as an IgG Fc fusion region peptide or leader or secretory sequence or a sequence which is employed for purification of the polypeptide or a proprotein sequence. Such fragments, derivatives and analogs are deemed to be within the scope of those skilled in the art from the teachings herein.

Of particular interest are substitutions of charged amino acids with another charged amino acid and with neutral or negatively charged amino acids. The latter results in proteins with reduced positive charge to improve the characteristics of a *M. jannaschii* ORF-encoded protein. The prevention of aggregation is highly desirable. Aggregation of proteins not only results in a loss of activity but can also be problematic when preparing pharmaceutical formulations, because they can be immunogenic. (Pinckard et al., Clin. Exp. Immunol. 2:331-340 (1967); Robbins et al., Diabetes 36:838-845 (1987); Cleland et al. Crit. Rev. Therapeutic Drug Carrier Systems 10:307-377 (1993)).

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As indicated, changes are preferably of a minor nature, such as conservative amino acid substitutions that do not significantly affect the folding or activity of the protein (see Table 1).

TABLE 1. Conservative Amino Acid Substitutions.

Aromatic	Phenylalanine Tryptophan	
Hydrophobic	Tyrosine Leucine Isoleucine Valine	
Polar	Glutamine Asparagine	
Basic	Arginine Lysine Histidine	
Acidic	Aspartic Acid Glutamic Acid	
Small	Alanine Serine Threonine	
	Methionine Glycine	

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Amino acids in a *M. jannaschii* ORF-encoded protein of the present invention that are essential for function can be identified by methods known in the art, such as site-directed mutagenesis or alanine-scanning mutagenesis

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(Cunningham and Wells, Science 244:1081-1085 (1989)). The latter procedure introduces single alanine mutations at every residue in the molecule.

The polypeptides of the present invention are preferably provided in an isolated form. By "isolated polypeptide" is intended a polypeptide removed from its native environment. Thus, a polypeptide produced and/or contained within a recombinant host cell is considered isolated for purposes of the present invention. Also intended as an "isolated polypeptide" are polypeptides that have been purified, partially or substantially, from a recombinant host cell. For example, a recombinantly produced version of a M. jannaschii ORF-encoded protein can be substantially purified by the one-step method described in Smith and Johnson, Gene 67:31-40 (1988).

The polypeptides of the present invention include the proteins encoded by (a) an ORF described in Table 2(a) or 3 or (b) an ORF described in Table 2(a) or 3, but minus the codon for the N-terminal methionine residue, if present, as well as polypeptides that have at least 90% similarity, more preferably at least 95% similarity, and still more preferably at least 96%, 97%, 98% or 99% similarity to a M. jannaschii ORF-encoded protein. Further polypeptides of the present invention include polypeptides at least 90% identical, more preferably at least 95% identical, still more preferably at least 96%, 97%, 98% or 99% identical to a M. jannaschii ORF-encoded protein.

By "% similarity" for two polypeptides is intended a similarity score produced by comparing the amino acid sequences of the two polypeptides using the Bestfit program (Wisconsin Sequence Analysis Package, Version 8 for Unix, Genetics Computer Group, University Research Park, 575 Science Drive, Madison, WI 53711) and the default settings for determining similarity. Bestfit uses the local homology algorithm of Smith and Waterman (Advances in Applied Mathematics 2:482-489, 1981) to find the best segment of similarity between two sequences.

By a polypeptide having an amino acid sequence at least, for example, 95% "identical" to a reference amino acid sequence of a *M. jannaschii* ORF-encoded protein is intended that the amino acid sequence of the polypeptide is

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identical to the reference sequence except that the polypeptide sequence may include up to five amino acid alterations per each 100 amino acids of the reference sequence. In other words, to obtain a polypeptide having an amino acid sequence at least 95% identical to a reference amino acid sequence, up to 5% of the amino acid residues in the reference sequence may be deleted or substituted with another amino acid, or a number of amino acids up to 5% of the total amino acid residues in the reference sequence may be inserted into the reference sequence. These alterations of the reference sequence may occur at the amino or carboxy terminal positions of the reference amino acid sequence or anywhere between those terminal positions, interspersed either individually among residues in the reference sequence or in one or more contiguous groups within the reference sequence.

As a practical matter, whether any particular polypeptide has an amino acid sequence at least 90%, 95%, 96%, 97%, 98% or 99% identical to the amino acid sequence of a *M. jannaschii* ORF-encoded protein can be determined conventionally using known computer programs such the Bestfit program (Wisconsin Sequence Analysis Package, Version 8 for Unix, Genetics Computer Group, University Research Park, 575 Science Drive, Madison, WI 53711). When using Bestfit or any other sequence alignment program to determine whether a particular sequence is, for instance, 95% identical to a reference sequence according to the present invention, the parameters are set, of course, such that the percentage of identity is calculated over the full length of the reference amino acid sequence and that gaps in homology of up to 5% of the total number of amino acid residues in the reference sequence are allowed.

As described in detail below, the polypeptides of the present invention can also be used to raise polyclonal and monoclonal antibodies, which are useful in assays for detecting *M. jannaschii* protein expression.

In another aspect, the invention provides a peptide or polypeptide comprising an epitope-bearing portion of a polypeptide of the invention. The epitope of this polypeptide portion is an immunogenic or antigenic epitope of a polypeptide of the invention. An "immunogenic epitope" is defined as a part of

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a protein that elicits an antibody response when the whole protein is the immunogen. These immunogenic epitopes are believed to be confined to a few loci on the molecule. On the other hand, a region of a protein molecule to which an antibody can bind is defined as an "antigenic epitope." The number of immunogenic epitopes of a protein generally is less than the number of antigenic epitopes. See, for instance, Geysen et al., Proc. Natl. Acad. Sci. USA 81:3998-4002 (1983).

As to the selection of peptides or polypeptides bearing an antigenic epitope (i.e., that contain a region of a protein molecule to which an antibody can bind), it is well known in that art that relatively short synthetic peptides that mimic part of a protein sequence are routinely capable of eliciting an antiserum that reacts with the partially mimicked protein. See, for instance, Sutcliffe, J. G., Shinnick, T. M., Green, N. and Learner, R.A. (1983). Antibodies that react with predetermined sites on proteins are described in Science 219:660-666. Peptides capable of eliciting protein-reactive sera are frequently represented in the primary sequence of a protein, can be characterized by a set of simple chemical rules, and are confined neither to immunodominant regions of intact proteins (i.e., immunogenic epitopes) nor to the amino or carboxyl terminals. Peptides that are extremely hydrophobic and those of six or fewer residues generally are ineffective at inducing antibodies that bind to the mimicked protein; longer, peptides, especially those containing proline residues, usually are effective. Sutcliffe et al., supra, at 661. For instance, 18 of 20 peptides designed according to these guidelines, containing 8-39 residues covering 75% of the sequence of the influenza virus hemagglutinin HA1 polypeptide chain, induced antibodies that reacted with the HA1 protein or intact virus; and 12/12 peptides from the MuLV polymerase and 18/18 from the rabies glycoprotein induced antibodies that precipitated the respective proteins.

Antigenic epitope-bearing peptides and polypeptides of the invention are therefore useful to raise antibodies, including monoclonal antibodies, that bind specifically to a polypeptide of the invention. Thus, a high proportion of hybridomas obtained by fusion of spleen cells from donors immunized with an

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antigen epitope-bearing peptide generally secrete antibody reactive with the native protein. Sutcliffe et al., supra, at 663. The antibodies raised by antigenic epitope-bearing peptides or polypeptides are useful to detect the mimicked protein, and antibodies to different peptides may be used for tracking the fate of various regions of a protein precursor which undergoes post-translational processing. The peptides and anti-peptide antibodies may be used in a variety of qualitative or quantitative assays for the mimicked protein, for instance in competition assays since it has been shown that even short peptides (e.g., about 9 amino acids) can bind and displace the larger peptides in immunoprecipitation assays. See, for instance, Wilson et al., Cell 37:767-778 (1984) at 777. The antipeptide antibodies of the invention also are useful for purification of the mimicked protein, for instance, by adsorption chromatography using methods well known in the art.

Antigenic epitope-bearing peptides and polypeptides of the invention designed according to the above guidelines preferably contain a sequence of at least seven, more preferably at least nine and most preferably between about 15 to about 30 amino acids contained within the amino acid sequence of a polypeptide of the invention. However, peptides or polypeptides comprising a larger portion of an amino acid sequence of a polypeptide of the invention, containing about 30 to about 50 amino acids, or any length up to and including the entire amino acid sequence of a polypeptide of the invention, also are considered epitope-bearing peptides or polypeptides of the invention and also are useful for inducing antibodies that react with the mimicked protein. Preferably, the amino acid sequence of the epitope-bearing peptide is selected to provide substantial solubility in aqueous solvents (i.e., the sequence includes relatively hydrophilic residues and highly hydrophobic sequences are preferably avoided); and sequences containing proline residues are particularly preferred.

The epitope-bearing peptides and polypeptides of the invention may be produced by any conventional means for making peptides or polypeptides including recombinant means using nucleic acid molecules of the invention. For instance, a short epitope-bearing amino acid sequence may be fused to a larger

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polypeptide which acts as a carrier during recombinant production and purification, as well as during immunization to produce anti-peptide antibodies. Epitope-bearing peptides also may be synthesized using known methods of chemical synthesis. For instance, Houghten has described a simple method for synthesis of large numbers of peptides, such as 10-20 mg of 248 different 13 residue peptides representing single amino acid variants of a segment of the HA1 polypeptide which were prepared and characterized (by ELISA-type binding studies) in less than four weeks. Houghten, R. A. (1985) General method for the rapid solid-phase synthesis of large numbers of peptides: specificity of antigen-antibody interaction at the level of individual amino acids. Proc. Natl. Acad. Sci. USA 82:5131-5135. This "Simultaneous Multiple Peptide Synthesis (SMPS)" process is further described in U.S. Patent No. 4,631,211 to Houghten et al. (1986). In this procedure the individual resins for the solid-phase synthesis of various peptides are contained in separate solvent-permeable packets, enabling the optimal use of the many identical repetitive steps involved in solid-phase methods. A completely manual procedure allows 500-1000 or more syntheses to be conducted simultaneously. Houghten et al., supra, at 5134.

Epitope-bearing peptides and polypeptides of the invention are used to induce antibodies according to methods well known in the art. See, for instance, Sutcliffe et al., supra; Wilson et al., supra; Chow, M. et al., Proc. Natl. Acad. Sci. USA 82:910-914; and Bittle, F. J. et al., J. Gen. Virol. 66:2347-2354 (1985). Generally, animals may be immunized with free peptide; however, anti-peptide antibody titer may be boosted by coupling of the peptide to a macromolecular carrier, such as keyhole limpet hemacyanin (KLH) or tetanus toxoid. For instance, peptides containing cysteine may be coupled to carrier using a linker such as m-maleimidobenzoyl-N-hydroxysuccinimide ester (MBS), while other peptides may be coupled to carrier using a more general linking agent such as glutaraldehyde. Animals such as rabbits, rats and mice are immunized with either free or carrier-coupled peptides, for instance, by intraperitoneal and/or intradermal injection of emulsions containing about 100 g peptide or carrier protein and Freund's adjuvant. Several booster injections may be needed, for

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instance, at intervals f about two weeks, to provide a useful titer of anti-peptide antibody which can be detected, for example, by ELISA assay using free peptide adsorbed to a solid surface. The titer of anti-peptide antibodies in serum from an immunized animal may be increased by selection of anti-peptide antibodies, for instance, by adsorption to the peptide on a solid support and elution of the selected antibodies according to methods well known in the art.

Immunogenic epitope-bearing peptides of the invention, i.e., those parts of a protein that elicit an antibody response when the whole protein is the immunogen, are identified according to methods known in the art. For instance, Geysen et al., supra, discloses a procedure for rapid concurrent synthesis on solid supports of hundreds of peptides of sufficient purity to react in an enzyme-linked immunosorbent assay. Interaction of synthesized peptides with antibodies is then easily detected without removing them from the support. In this manner a peptide bearing an immunogenic epitope of a desired protein may be identified routinely by one of ordinary skill in the art. For instance, the immunologically important epitope in the coat protein of foot-and-mouth disease virus was located by Geysen et al. with a resolution of seven amino acids by synthesis of an overlapping set of all 208 possible hexapeptides covering the entire 213 amino acid sequence of the protein. Then, a complete replacement set of peptides in which all 20 amino acids were substituted in turn at every position within the epitope were synthesized, and the particular amino acids conferring specificity for the reaction with antibody were determined. Thus, peptide analogs of the epitope-bearing peptides of the invention can be made routinely by this method. U.S. Patent No. 4,708,781 to Geysen (1987) further describes this method of identifying a peptide bearing an immunogenic epitope of a desired protein.

Further still, U.S. Patent No. 5,194,392 to Geysen (1990) describes a general method of detecting or determining the sequence of monomers (amino acids or other compounds) which is a topological equivalent of the epitope (i.e., a "mimotope") which is complementary to a particular paratope (antigen binding site) of an antibody of interest. More generally, U.S. Patent No. 4,433,092 to Geysen (1989) describes a method of detecting or determining a sequence of

monomers which is a topographical equivalent f a ligand which is complementary to the ligand binding site f a particular receptor of interest. Similarly, U.S. Patent No. 5,480,971 to Houghten, R. A. et al. (1996) on Peralkylated Oligopeptide Mixtures discloses linear C₁-C₇-alkyl peralkylated oligopeptides and sets and libraries of such peptides, as well as methods for using such oligopeptide sets and libraries for determining the sequence of a peralkylated oligopeptide that preferentially binds to an acceptor molecule of interest. Thus, non-peptide analogs of the epitope-bearing peptides of the invention also can be made routinely by these methods.

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The entire disclosure of each document cited in this section on "Polypeptides and Peptides" is hereby incorporated herein by reference.

As one of skill in the art will appreciate, the polypeptides of the present invention and the epitope-bearing fragments thereof described above can be combined with parts of the constant domain of immunoglobulins (IgG), resulting in chimeric polypeptides. These fusion proteins facilitate purification and show an increased half-life *in vivo*. This has been demonstrated, e.g., for chimeric proteins consisting of the first two domains of the human CD4-polypeptide and various domains of the constant regions of the heavy or light chains of mammalian immunoglobulins (EPA 394,827; Traunecker *et al.*, *Nature 331*:84-86 (1988)). Fusion proteins that have a disulfide-linked dimeric structure due to the IgG part can also be more efficient in binding and neutralizing other molecules than the monomeric protein or protein fragment alone (Fountoulakis *et al.*, *J Biochem 270*:3958-3964 (1995)).

Protein Function

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Each ORF described in Table 2(a) was assigned to biological role categories adapted from Riley, M., Microbiology Reviews 57(4):862 (1993)). This allows the skilled artisan to determine a function for each identified coding sequence. For example, a partial list of the M. jannaschii protein functions provided in Table 2(a) includes: methanogenesis, amino acid biosynthesis, cell

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division, detoxificati n, protein secreti n, transformati n, central intermediary metabolism, energy metabolism, degradation of DNA, DNA replication, restriction, modification, recombination and repair, transcription, RNA processing, translation, degradation of proteins, peptides and glycopeptides, ribosomal proteins, translation factors, transport, tRNA modification, and drug and analog sensitivity. A more detailed description of several of these functions is provided in Example 1 below.

Diagnostic Assays

The present invention further provides methods to identify the expression of an ORF of the present invention, or homolog thereof, in a test sample, using one of the DFs or antibodies of the present invention. Such methods involve incubating a test sample with one or more of the antibodies or one or more of the DFs of the present invention and assaying for binding of the DFs or antibodies to components within the test sample.

Conditions for incubating a DF or antibody with a test sample vary. Incubation conditions depend on the format employed in the assay, the detection methods employed, and the type and nature of the DF or antibody used in the assay. One skilled in the art will recognize that any one of the commonly available hybridization, amplification or immunological assay formats can readily be adapted to employ the DFs or antibodies of the present invention. Examples of such assays can be found in Chard, T., An Introduction to Radioimmunoassay and Related Techniques, Elsevier Science Publishers, Amsterdam, The Netherlands (1986); Bullock, G.R. et al., Techniques in Immunocytochemistry, Academic Press, Orlando, FL Vol. 1 (1982), Vol. 2 (1983), Vol. 3 (1985); Tijssen, P., Practice and Theory of Enzyme Immunoassays: Laboratory Techniques in Biochemistry and Molecular Biology, Elsevier Science Publishers, Amsterdam, The Netherlands (1985).

The test samples of the present invention include cells, protein or membrane extracts of cells. The test sample used in the above-described method

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will vary based on the assay format, nature of the detection method and the cells or extracts used as the sample to be assayed. Methods for preparing protein extracts or membrane extracts of cells are well known in the art and can be readily be adapted in order to obtain a sample which is compatible with the system utilized.

In another embodiment of the present invention, kits are provided which contain the necessary reagents to carry out the assays of the present invention. Specifically, the invention provides a compartmentalized kit to receive, in close confinement, one or more containers including comprising: (a) a first container comprising one of the DFs or antibodies of the present invention; and (b) one or more other containers comprising one or more of the following: wash reagents, reagents capable of detecting presence of a bound DF or antibody.

A compartmentalized kit includes any kit in which reagents are contained in separate containers. Such containers include small glass containers, plastic containers or strips of plastic or paper. Such containers allow one to efficiently transfer reagents from one compartment to another compartment such that the samples and reagents are not cross-contaminated, and the agents or solutions of each container can be added in a quantitative fashion from one compartment to another. Such containers will include a container which will accept the test sample, a container which contains the antibodies used in the assay, containers which contain wash reagents (such as phosphate buffered saline, Tris-buffers, etc.), and containers which contain the reagents used to detect the bound antibody or DF.

Types of detection reagents include labeled nucleic acid probes, labeled secondary antibodies, or in the alternative, if the primary antibody is labeled, the enzymatic, or antibody binding reagents that are capable of reacting with the labeled antibody. One skilled in the art will readily recognize that the disclosed DFs and antibodies of the present invention can be readily incorporated into one of the established kit formats that are well known in the art.

Screening Assay for Binding Agents

Using the isolated proteins described herein, the present invention further provides methods of obtaining and identifying agents that bind to a protein encoded by a M. jannaschii ORF or to a fragment thereof.

The method involves:

- (a) contacting an agent with an isolated protein encoded by a M. jannaschii ORF, or an isolated fragment thereof; and
- (b) determining whether the agent binds to said protein or said fragment.

The agents screened in the above assay can be, but are not limited to, peptides, carbohydrates, vitamin derivatives, or other pharmaceutical agents. The agents can be selected and screened at random or rationally selected or designed using protein modeling techniques. For random screening, agents such as peptides, carbohydrates, pharmaceutical agents and the like are selected at random and are assayed for their ability to bind to the protein encoded by an ORF of the present invention.

Alternatively, agents may be rationally selected or designed. As used herein, an agent is said to be "rationally selected or designed" when the agent is chosen based on the configuration of the particular protein. For example, one skilled in the art can readily adapt currently available procedures to generate peptides, pharmaceutical agents and the like capable of binding to a specific peptide sequence in order to generate rationally designed antipeptide peptides, for example see Hurby et al., Application of Synthetic Peptides: Antisense Peptides, In Synthetic Peptides, A User's Guide, W.H. Freeman, NY (1992), pp. 289-307, and Kaspczak et al., Biochemistry 28:9230-8 (1989), or pharmaceutical agents, or the like.

In addition to the foregoing, one class of agents of the present invention, can be used to control gene expression through binding to one of the ORFs or EMFs of the present invention. As described above, such agents can be randomly

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screened r rationally designed and selected. Targeting the ORF or EMF all ws a skilled artisan to design sequence specific r element specific agents, modulating the expression of either a single ORF or multiple ORFs that rely on the same EMF for expression control.

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One class of DNA binding agents are those that contain nucleotide base residues that hybridize or form a triple helix by binding to DNA or RNA. Such agents can be based on the classic phosphodiester, ribonucleic acid backbone, or can be a variety of sulfhydryl or polymeric derivatives having base attachment capacity.

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Agents suitable for use in these methods usually contain 20 to 40 bases and are designed to be complementary to a region of the gene involved in transcription (triple helix - see Lee et al., Nucl. Acids Res. 6:3073 (1979); Cooney et al., Science 241:456 (1988); and Dervan et al., Science 251: 1360 (1991)) or to the mRNA itself (antisense - Okano, J. Neurochem. 56:560 (1991); Oligodeoxynucleotides as Antisense Inhibitors of Gene Expression, CRC Press, Boca Raton, FL (1988)). Triple helix-formation optimally results in a shut-off of RNA transcription from DNA, while antisense RNA hybridization blocks translation of an mRNA molecule into polypeptide. Both techniques have been demonstrated to be effective in model systems. Information contained in the sequences of the present invention is necessary for the design of an antisense or triple helix oligonucleotide and other DNA binding agents.

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Computer Related Embodiments

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The nucleotide sequence provided in SEQ ID NO:1, 2, or 3, a representative fragment thereof, or a nucleotide sequence at least 99.9% identical to the sequence provided in SEQ ID NO:1, 2, or 3, can be "provided" in a variety of mediums to facilitate use thereof. As used herein, provided refers to a manufacture, other than an isolated nucleic acid molecule, that contains a nucleotide sequence of the present invention, i.e., the nucleotide sequence provided in SEQ ID NO:1, 2, or 3, a representative fragment thereof, or a

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nucleotide sequence at least 99.9% identical to SEQ ID NO:1, 2, or 3. Such a manufacture provides the *M. jannaschii* genome or a subset thereof (e.g., a *M. jannaschii* open reading frame (ORF)) in a form that allows a skilled artisan to examine the manufacture using means not directly applicable to examining the *M. jannaschii* genome or a subset thereof as it exists in nature or in purified form.

In one application of this embodiment, a nucleotide sequence of the present invention can be recorded on computer readable media. As used herein, "computer readable media" refers to any medium that can be read and accessed directly by a computer. Such media include, but are not limited to: magnetic storage media, such as floppy discs, hard disc storage medium, and magnetic tape; optical storage media such as CD-ROM; electrical storage media such as RAM and ROM; and hybrids of these categories such as magnetic/optical storage media. A skilled artisan can readily appreciate how any of the presently known computer readable mediums can be used to create a manufacture comprising computer readable medium having recorded thereon a nucleotide sequence of the present invention.

As used herein, "recorded" refers to a process for storing information on computer readable medium. A skilled artisan can readily adopt any of the presently know methods for recording information on computer readable medium to generate manufactures comprising the nucleotide sequence information of the present invention. A variety of data storage structures are available to a skilled artisan for creating a computer readable medium having recorded thereon a nucleotide sequence of the present invention. The choice of the data storage structure will generally be based on the means chosen to access the stored information. In addition, a variety of data processor programs and formats can be used to store the nucleotide sequence information of the present invention on computer readable medium. The sequence information can be represented in a word processing text file, formatted in commercially-available software such as WordPerfect and MicroSoft Word, or represented in the form of an ASCII file, stored in a database application, such as DB2, Sybase, Oracle, or the like. A skilled artisan can readily adapt any number of dataprocessor structuring formats

(e.g. text file or database) in order to obtain computer readable medium having recorded thereon the nucleotide sequence information of the present invention.

By providing the nucleotide sequence of SEQ ID NO:1, 2, or 3, a representative fragment thereof, or a nucleotide sequence at least 99.9% identical to SEQ ID NO:1, 2, or 3, in computer readable form, a skilled artisan can routinely access the sequence information for a variety of purposes. Computer software is publicly available which allows a skilled artisan to access sequence information provided in a computer readable medium. The examples which follow demonstrate how software which implements the BLAST (Altschul et al., J. Mol. Biol. 215:403-410 (1990)) and BLAZE (Brutlag et al., Comp. Chem. 17:203-207 (1993)) search algorithms on a Sybase system can be used to identify open reading frames (ORFs) within the M. jannaschii genome that contain homology to ORFs or proteins from other organisms. Such ORFs are proteinencoding fragments within the M. jannaschii genome and are useful in producing commercially important proteins such as enzymes used in methanogenesis, amino acid biosynthesis, metabolism, fermentation, transcription, translation, RNA processing, nucleic acid and protein degradation, protein modification, and DNA replication, restriction, modification, recombination, and repair. comprehensive list of ORFs encoding commercially important M. jannaschii proteins is provided in Tables 2(a) and 3.

The present invention further provides systems, particularly computer-based systems, which contain the sequence information described herein. Such systems are designed to identify commercially important fragments of the *M. jannaschii* genome. As used herein, "a computer-based system" refers to the hardware means, software means, and data storage means used to analyze the nucleotide sequence information of the present invention. The minimum hardware means of the computer-based systems of the present invention comprises a central processing unit (CPU), input means, output means, and data storage means. A skilled artisan can readily appreciate that any one of the currently available computer-based system are suitable for use in the present invention.

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As indicated above, the computer-based systems f the present invention comprise a data storage means having stored therein a nucleotide sequence of the present invention and the necessary hardware means and software means for supporting and implementing a search means. As used herein, "data storage means" refers to memory that can store nucleotide sequence information of the present invention, or a memory access means which can access manufactures having recorded thereon the nucleotide sequence information of the present invention. As used herein, "search means" refers to one or more programs which are implemented on the computer-based system to compare a target sequence or target structural motif with the sequence information stored within the data storage means. Search means are used to identify fragments or regions of the M. jannaschii genome that match a particular target sequence or target motif. A variety of known algorithms are disclosed publicly and a variety of commercially available software for conducting search means are available and can be used in the computer-based systems of the present invention. Examples of such software include, but are not limited to, MacPattern (EMBL), BLASTN and BLASTX (NCBIA). A skilled artisan can readily recognize that any one of the available algorithms or implementing software packages for conducting homology searches can be adapted for use in the present computer-based systems.

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As used herein, a "target sequence" can be any DNA or amino acid sequence of six or more nucleotides or two or more amino acids. A skilled artisan can readily recognize that the longer a target sequence is, the less likely a target sequence will be present as a random occurrence in the database. The most preferred sequence length of a target sequence is from about 10 to 100 amino acids or from about 30 to 300 nucleotide residues. However, it is well recognized that during searches for commercially important fragments of the M. jannaschii genome, such as sequence fragments involved in gene expression and protein processing, may be of shorter length.

As used herein, "a target structural motif," or "target motif," refers to any rationally selected sequence r combination of sequences in which the sequence(s) are chosen based on a three-dimensional configuration which is

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formed upon the folding of the target motif. There are a variety f target motifs known in the art. Protein target motifs include, but are not limited to, enzymic active sites and signal sequences. Nucleic acid target motifs include, but are not limited to, promoter sequences, hairpin structures and inducible expression elements (protein binding sequences).

Thus, the present invention further provides an input means for receiving a target sequence, a data storage means for storing the target sequence and the homologous *M. jannaschii* sequence identified using a search means as described above, and an output means for outputting the identified homologous *M. jannaschii* sequence. A variety of structural formats for the input and output means can be used to input and output information in the computer-based systems of the present invention. A preferred format for an output means ranks fragments of the *M. jannaschii* genome possessing varying degrees of homology to the target sequence or target motif. Such presentation provides a skilled artisan with a ranking of sequences which contain various amounts of the target sequence or target motif and identifies the degree of homology contained in the identified fragment.

A variety of comparing means can be used to compare a target sequence or target motif with the data storage means to identify sequence fragments of the *M. jannaschii* genome. For example, implementing software which implement the BLAST and BLAZE algorithms (Altschul *et al.*, *J. Mol. Biol. 215*:403-410 (1990)) can be used to identify open reading frames within the *M. jannaschii* genome. A skilled artisan can readily recognize that any one of the publicly available homology search programs can be used as the search means for the computer-based systems of the present invention.

One application of this embodiment is provided in Figure 4. Figure 4 provides a block diagram of a computer system 102 that can be used to implement the present invention. The computer system 102 includes a processor 106 connected to a bus 104. Also connected to the bus 104 are a main memory 108 (preferably implemented as random access memory, RAM) and a variety of secondary storage devices 110, such as a hard drive 112 and a removable medium

storage device 114. The removable medium storage device 114 may represent, for example, a floppy disk drive, a CD-ROM drive, a magnetic tape drive, etc. A removable storage medium 116 (such as a floppy disk, a compact disk, a magnetic tape, etc.) containing control logic and/or data recorded therein may be inserted into the removable medium storage device 114. The computer system 102 includes appropriate software for reading the control logic and/or the data from the removable medium storage device 114 once inserted in the removable medium storage device 114.

A nucleotide sequence of the present invention may be stored in a well known manner in the main memory 108, any of the secondary storage devices 110, and/or a removable storage medium 116. Software for accessing and processing the genomic sequence (such as search tools, comparing tools, etc.) reside in main memory 108 during execution.

Having generally described the invention, the same will be more readily understood by reference to the following examples, which are provided by way of illustration and are not intended as limiting.

Experimental

Complete genome sequence of the methanogenic archaeon, Methanococcus jannaschii

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A whole genome random sequencing method (Fleischmann, R.D., et al., Science 269:496 (1995); Fraser, C.M., et al., Science 270:397 (1995)) was used to obtain the complete genome sequence for M. jannaschii. A small insert plasmid library (2.5 Kbp average insert size) and a large insert lambda library (16 Kbp average insert size) were used as substrates for sequencing. The lambda library was used to form a genome scaffold and to verify the orientation and integrity of the contigs formed from the assembly of sequences from the plasmid library. All clones were sequenced from both ends to aid in redering of contigs during the sequence assembly process. The average length of sequencing reads was 481 bp. A total of 36,718 sequences were assembled by means of the TIGR

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Assembler (Fleischmann, R.D., et al., Science 269:496 (1995); Fraser, C.M., et al., Science 270:397 (1995); Sutton G., et al., Genome Sci. Tech. 1:9 (1995)). Sequence and physical gaps were closed using a combination of strategies (Fleischmann, R.D., et al., Science 269:496 (1995); Fraser, C.M., et al., Science 270:397 (1995)). The colinearity of the in vivo genome to the genome sequence was confirmed by comparing restriction fragments from six. rare cutter, restriction enzymes (Aat II, BamHI, Bgl II, Kpn I, Sma I, and Sst II) to those predicted from the sequence data. Additional confidence in the colinearity was provided by the genome scaffold produced by sequence pairs from 339 largeinsert lambda clones, which covered 88% of the main chromosome. Open reading frames (ORFs) and predicted protein-coding regions were identified as described (Fleischmann, R.D., et al., Science 269:496 (1995); Fraser, C.M., et al., Science 270:397 (1995)) with some modification. In particular, the statistical prediction of M. jannaschii genes was performed with GeneMark (Borodovsky, M. & McIninch, J. Comput. Chem. 17:123 (1993)). Regular GeneMark uses nonhomogeneous Markov models derived from a training set of coding sequences and ordinary Markov models derived from a training set of noncoding sequences. Only a single 16S ribosomal RNA sequence of M. jannaschii was available in the public sequence databases before the whole genome sequence described here. Thus, the initial training set to determine parameters of a coding sequence Markov model was chosen as a set of ORFs > 1000 nucleotides (nt). As an initial model for non-coding sequences, a zero-order Markov model with genomespecific nucleotide frequencies was used. The initial models were used at the first prediction step. The results of the first prediction were then used to compile a set of putative genes used at the second training step. Alternate rounds of training and predicting were continued until the set of predicted genes stabilized and the parameters of the final fourth-order model of coding sequences were derived. The regions predicted as noncoding were then used as a training set for a final model for noncoding regions. Cross-validation simulations demonstrated that the GeneMark program trained as described above was able to correctly identify coding regions of at least 96 nt in 94% of the cases and noncoding regions of the

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same length in 96% of the cases. These values assume that the self-training method produced correct sequence annotation for compiled control sets. Comparison with the results obtained by searches against a nonredundant protein database (Fleischmann, R.D., et al., Science 269:496 (1995); Fraser, C.M., et al., Science 270:397 (1995)) demonstrated that almost all genes identified by sequence similarity were predicted by the GeneMark program as well. This observation provides additional confidence in genes predicted by GeneMark whose protein translations did not show significant similarity to known protein sequences. The predicted protein-coding regions were search against the Blocks database (Henikoff, S. & Henikoff, J.G., Genomics 19:97 (1994)] by means of BLIMPS (Wallace, J.C. & Henikoff, S., CABIOS 8:249 (1992)) to verify putative identifications and to identify potential functional motifs in predicted proteincoding regions that had no database match. Genes were assigned to known metabolic pathways. When a gene appeared to be missing from a pathway, the unassigned ORFs and the complete M. jannaschii genome sequence were searched with specific query sequences or motifs from the Blocks database. Hydrophobicity plots were performed on all predicted protein-coding regions by means of the Kyte-Doolittle algorithm (Kyte, J. & Doolittle, R.F., J. Mol. Biol. 157:105 (1982)) to identify potentially functionally relevant signatures in these sequences.

The *M. jannaschii* genome comprises three physically distinct elements: i) a large circular chromosome of 1,664,976 base pairs (bp) (SEQ ID NO:1), which contains 1682 predicted protein-coding regions and has a G+C content of 31.4%; ii) a large circular extrachromosomal element (ECE) (Zhao, H., *et al.*, *Arch. Microbiol. 150*:178 (1988)) of 58,407 bp (SEQ ID NO:2), which contains 44 predicted protein coding regions and has a G+C content of 28.2%; and iii) a small circular ECE (Zhao, H., *et al.*, *Arch. Microbiol. 150*:178 (1988)) of 16,550 bp (SEQ ID NO:3), which contains 12 predicted protein coding regions, and has a G+C content of 28.8%. With respect to its shape, size, G+C content, and gene density the main chromosome resembles that of *H. influenzae*. However, here the resemblance stops.

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Of the 1743 predicted protein-coding regi ns reported previously for H. influenzae, 78% had a match in the public sequence database (Fleischmann, R.D., et al., Science 269:496 (1995); Frascr, C.M., et al., Science 270:397 (1995)). Of these, 58% were matches to genes with reasonably well defined function, while 20% were matches to genes whose function was undefined. Similar observations were made for the M. genitalium genome (Fleischmann, R.D., et al., Science 269:496 (1995); Fraser, C.M., et al., Science 270:397 (1995)). Eighty-three percent of the predicted protein coding regions from M. genitalium have a counterpart in the H. influenzae genome. In contrast, only 38% of the predicted protein-coding regions from M. jannaschii match a gene in the database that could be assigned a putative cellular role with high confidence; 6% of the predicted protein-coding regions had matches to hypothetical proteins (Tables 2-3). Approximately 100 genes in M. jannaschii had marginal similarity to genes or segments of genes from the public sequence databases and could not be assigned a putative cellular role with high confidence. Only 11% of the predicted protein-coding regions from H. influenzae and 17% of the predicted protein coding regions from M. genitalium matched a predicted protein coding region from M. jannaschii. Clearly the M. jannaschii genome, and undoubtedly, therefore, all archaeal genomes are remarkably unique, as the phylogenetic position of these organisms would suggest.

Energy production in *M. jannaschii* occurs via the reduction of CO₂ with H₂ to produce methane. Genes for all of the known enzymes and enzyme complexes associated with methanogenesis (DiMarco, A.A., et al., Ann. Rev. Biochem. 59:355 (1990)) were identified in *M. jannaschii*, the sequence and order of which are typical of methanogens. *M. jannaschii* appears to use both H₂ and formate as substrates for methanogenesis, but lacks the genes to use methanol or acetate. The ability to fix nitrogen has been demonstrated in a number of methanogens (Belay, N., et al., Nature 312:286 (1984)) and all of the genes necessary for this pathway have been identified in *M. jannaschii* (Tables 2-3). In addition to its anabolic pathways, several scavenging molecules have been

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identified in *M. jannaschii* that probably play a role in importing small organic compounds, such as amino acids, from the environment (Tables 2-3).

Three different pathways are known for the fixation of CO₂ into organic carbon: the non-cyclic, reductive acetyl-coenzyme A-carbon monoxide dehydrogenase pathway (Ljungdahl-Wood pathway), the reductive trichloroacetic acid (TCA) cycle, and the Calvin cycle. Methanogens fix carbon by the Ljungdahl-Wood pathway (Wood, H.G., et al., TIBS 11:14 (1986)), which is facilitated by the carbon monoxide dehydrogenease enzyme complex (CODH) (Blaat, M., Antonie van Leewenhoek 66:187 (1994)). The complete Ljungdahl-Wood pathway, encoded in the M. jannaschii genome, depends on the methyl carbon in methanogenesis; however, methanogenesis can occur independently of carbon fixation.

Although genes encoding two enzymes required for gluconeogenesis (glucopyruvate oxidoreductase and phosphoenolpyruvate synthase) were found in the *M. jannaschii* genome, genes encoding other key intermediates of gluconeogenesis (fructose bisphosphatase and fructose 1,6-bisphosphate aldolase) were not been identified. Glucose catabolism by glycolysis also requires the aldolase, as well as phosphofructokinase, an enzyme that also was not found in *M. jannaschii* and has not been detected in any of the Archaea. In addition, genes specific for the Entner-Doudoroff pathway, an alternative pathway used by some microbes for the catabolism of glucose, were not identified in the genomic sequence. The presence of a number of nearly complete metabolic pathways suggests that some key genes are not recognizable at the sequence level, although we cannot exclude the possibility that *M. jannaschii* may use alternative metabolic pathways.

In general, M. jannaschii genes that encode proteins involved in the transport of small inorganic ions into the cell are homologs of bacterial genes. The genome includes many representatives of the ABC transporter family, as well as genes for exporting heavy metals (e.g., the chromate-resistance protein) and other toxic compounds (e.g., the norA drug efflux pump locus).

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M re than 20 predicted protein-coding regions have sequence similarity to polysaccharide biosynthetic enzymes. These genes have nly bacterial homologs or are most closely related to their bacterial counterparts. The identified polysaccharide biosynthetic genes in M. jannaschii include those for the interconversion of sugars, activation of sugars to nucleotide sugars, and glycosyltransferases for the polymerization of nucleotide sugars into oligo- and polysaccharides that are subsequently incorporated into surface structures (Hartmann, E. and König, H., Arch. Microbiol. 151:274 (1989)). In an arrangement reminiscent of bacterial polysaccharide biosynthesis genes, many of the genes for M. jannaschii polysaccharide production are clustered together (Tables 2-3). The G+C content in this region is <95% of that in the rest of the M. jannaschii genome. A similar observation was made in Salmonella typhimurium (Jiang, X.M., et al., Mol. Microbiol. 5:695 (1991)) in which the gene cluster for lipopolysaccharide O antigen has a significantly lower G+C ratio than the rest of the genome. In that case, the difference in G+C content was interpreted as meaning that the region originated by lateral transfer from another organism.

Of the three main multicomponent information processing systems (transcription, translation, and replication), translation appears the most universal in its overall makeup in that the basic translation machinery is similar in all three domains of life. *M. jannaschii* has two ribosomal RNA operons, designated A and B, and a separate 5S RNA gene that is associated with several transfer RNAs (tRNAs). Operon A has the organization, 16S - 23S - 5S, whereas operon B lacks the 5S component. An alanine tRNA is situated in the spacer region between the 16S and 23S subunits in both operons. The majority of proteins associated with the ribosomal subunits (especially the small subunit) are present in both Bacteria and Eukaryotes. However, the relatively protein-rich eukaryotic ribosome contains additional ribosomal proteins not found in the bacterial ribosome. A smaller number of bacteria-specific ribosomal proteins exist as well. The *M. jannaschii* genome contains all ribosomal proteins that are common to eukaryotes and bacteria. It shows no homologs of the bacterial-specific ribosomal proteins, but does possess homologs of a number of the eukaryotic-specific ones.

Homologs of all archaea-specific ribosomal proteins that have been reported to date (Lechner, K., et al., J. Mol. Evol. 29:20 (1989); Köpke, A.K.E. and Wittmann-Liebold, B., Can. J. Microbiol. 35:11 (1989)) are found in M. jannaschii.

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Thirty-seven tRNA genes were identified in the *M. jannaschii* genome. Almost all amino acids encoded by two codons have a single tRNA, except for glutamic acid, which has two. Both an initiator and an internal methionyl tRNA are present. The two pyrimidine-ending isoleucine codons are covered by a single tRNA, while the third (AUA) seems covered by a related tRNA having a CAU anticodon. A single tRNA appears to cover the three isoleucine codons. Those amino acids encoded by four codons each have two tRNAs, one to cover the Y-, the other the R-ending, codons. Valine has a third tRNA, which is

As previously shown for other archaea (Iwabe, N., et al., Proc. Natl. Acad. Sci. USA 86:9355 (1989); Gogarten J.P., et al., Proc. Natl. Acad. Sci. USA 86:6661 (1989); Brown, J.R. and Doolittle, W.F., Proc. Natl. Acad. Sci. USA 92:2441 (1995)), the Methanococcus translation elongation factors EF-1 α (EF-Tu in bacteria) and EF-2 (EF-G in bacteria) are most similar to their eukaryotic counterparts. In addition, the M. jannaschii genome contains 11 translation initiation factor genes. Three of these genes encode the subunits homologous to those of the eukaryotic IF-2, and are reported here in the Archaea for the first time. A fourth initiation factor gene that encodes a second IF-2 is also found in M. jannaschii. This additional IF-2 gene is most closely related to the yeast protein FUN12 which, in turn, appears to be a homolog of the bacterial IF-2. It is not known which of the two IF-2-like initiation factors identified in M. jannaschii plays a role in directing the initiator tRNA to the start site of the mRNA. The fifth identified initiation factor gene in M. jannaschii encodes IF-1A, which has no bacterial homolog. The sixth gene encodes the hypusinecontaining initiation factor eIF-5a. Two subunits of the translation initiation factor eIF-2B were identified in M. jannaschii. Finally, three putative adenososine 5'-triphosphate (ATP)-dependent helicases were identified that belong to the eIF-4a family of translation initiation factors.

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specific for the GUG codon; and alanine has three tRNAs (two of which are in the spacer regions separating the 16S and 23S subunits in the two ribosomal RNA operons). Leucine, serine and arginine, all of which have six codons, each posses three corresponding tRNAs. The genes for the internal methionine and tryptophan tRNAs contain introns in the region of their anti-codon loops.

A tRNA also exists for selenocysteine (UGA codon). At least four genes in M. jannaschii contain internal stop codons that are potential selenocysteine codons: the α chain of formate dehydrogenase, coenzyme F420 reducing hydrogenase, β -chain tungsten formyl methanofuran dehydrogenase, and a heterodisulfide reductase. Three genes with a putative role in selenocysteine metabolism were identified by their similarity to the sel genes from other organisms (Tables 2-3).

Recognizable homologs for four of the aminoacyl-tRNA synthetases (glutamine, asparagine, lysine, and cysteine) were not identified in the M. jannaschii genome. The absence of a glutaminyl-tRNA synthetase is not surprising in that a number of organisms, including at least one archaeon, have none (Wilcox, M., Eur. J. Biochem. 11:405 (1969); Martin, N.C., et al., J. Mol. Biol. 101:285 (1976); Martin, N.C., et al., Biochemistry 16:4672 (1977); Schon, A., et al., Biochimie 70:391 (1988); Soll, D. and RajBhandary, U., Eds. Am. Soc. for Microbiol. (1995)). In these instances, glutaminyl tRNA charging involves a post-charging conversion mechanism whereby the tRNA is charged by the glutamyl-tRNA synthetase with glutamic acid, which then is enzymatically converted to glutamine. A post-charging conversion is also involved in selenocysteine charging via the seryl-tRNA synthetase. A similar mechanism has been proposed for asparagine charging, but has never been demonstrated (Wilcox, M., Eur. J. Biochem. 11:405 (1969); Martin, N.C., et al., J. Mol. Biol. 101:285 (1976); Martin, N.C., et al., Biochemistry 16:4672 (1977); Schon, A., et al., Biochimie 70:391 (1988); Soll, D. and RajBhandary, U., Eds. Am. Soc. for Microbiol. (1995)). The inability to find homologs of the lysine and cysteine aminoacyl-tRNA synthetases is surprising because bacterial and eukaryotic versions in each instance show clear homology.

Aminoacyl-tRNA synthetases of *M. jannaschii* and ther archaea resemble eukaryotic synthetases more closely than they resemble bacterial forms. The tryptophanyl synthetase is one of the more notable examples, because the *M. jannaschii* and eukaryotic version do not appear to be specifically related to the bacterial version (de Pouplana, R., et al., Proc. Natl. Acad. Sci., USA 93:166 (1996)). Two versions of the glycyl synthetase are known in bacteria, one that is very unlike the version found in Archaea and Eukaryote and one that is an obvious homolog of it (Wagner, E.A., et al., J. Bacteriol: 177:5179 (1995); Logan, D.T., et al., EMBO J. 14:4156 (1995)).

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Eleven genes encoding subunits of the DNA-dependent RNA polymerase were identified in the *M. jannaschii* genome. The sequence similarity between the subunits and their homologs in *Sulfolobus acidocaldarius* supports the evolutionary unity of the archaeal polymerase complex (Woese, C.R. and Wolfe, R.S., Eds. *The Bacteria, vol. VIII* (Academic Press, NY, 1985); Langer, D., et al., *Proc. Natl. Acad. Sci. 92:5768* (1995); Lanzendoerfer, M. et al., *System. Appl. Microbiol. 16:656* (1994)). All of the subunits found in *M. jannaschii* show greater similarity to their eukaryotic counterparts than to the bacterial homologs. The genes encoding the five largest subunits (A', A'', B', B'', D) have homologs in all organisms. Six genes encode subunits shared only by Archaea and Eukaryotes (E, H, K, L, and N). The *M. jannaschii* homolog of the *S. acidocaldarius* subunit E is split into two genes designated E' and E''. *Sulfolobus acidocaldarius* also contains two additional small subunits of RNA polymerase, designated G and F, that have no counterparts in either Bacteria or Eukaryotes. No homolog of these subunits was identified in *M. jannaschii*.

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The archaeal transcription initiation system is essentially the same as that found in Eukaryotes, and is radically different from the bacterial version (Klenk, H.P. and Doolittle, W.F., Curr. Biol. 4:920 (1994)). The central molecules in the former systems are the TATA-binding protein (TBP) and transcription factor B (TFIIB and TFIIIB in Eukaryotes, or simply TFB). In the eukaryotic systems, TBP and TFB are parts of larger complexes, and additional factors (such as

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TFIIA and TFIIF) are used in the transcription process. However, the M. jannaschii genome does not contain obvious homologs of TFIIA and TFIIF.

Several components of the replication machinery were identified in M jannaschii. The M jannaschii genome appears to encode a single DNA-dependent polymerase that is a member of the B family of polymerases (Bernard, A., et al., EMBO J. 6:4219 (1987); Cullman, G., et al., Molec. Cell Biol. 15:4661 (1995); Uemori, T., et al., J. Bacteriol. 117:2164 (1995); Delarue, M., et al., Prot. Engineer. 3:461 (1990); Gavin, K.A., et al., Science 270:1667 (1995)). The polymerase shares sequence similarity and three motifs with other family B polymerases, including eukaryotic α , γ , and ϵ polymerases, bacterial polymerase II, and several archaeal polymerases. However, it is not homologous to bacterial polymerase I and has no homologs in H. influenzae or M. genitalium.

Primer recognition by the polymerase takes place through a structurespecific DNA binding complex, the replication factor complex (rfc) (Bernard, A., et al., EMBO J. 6:4219 (1987); Cullman, G., et al., Molec. Cell Biol. 15:4661 (1995); Uemori, T., et al., J. Bacteriol. 117:2164 (1995); Delarue, M., et al., Prot. Engineer. 3:461 (1990); Gavin, K.A., et al., Science 270:1667 (1995)). In humans and yeast, the rfc is composed of five proteins: a large subunit and four small subunits that have an associated adenosine triphosphatase (ATPase) activity stimulated by proliferating cell nuclear antigen (PCNA). Two genes in M. jannaschii are putative members of a eukaryotic-like replication factor complex. One of the genes in M. jannaschii is a putative homolog of the large subunit of the rfc, whereas the second is a putative homolog of one of the small subunits. Among Eukaryotes, the rfc proteins share sequence similarity in eight signature domains (Bernard, A., et al., EMBO J. 6:4219 (1987); Cullman, G., et al., Molec. Cell Biol. 15:4661 (1995); Uemori, T., et al., J. Bacteriol. 117:2164 (1995); Delarue, M., et al., Prot. Engineer. 3:461 (1990); Gavin, K.A., et al., Science 270:1667 (1995)). Domain I is conserved only in the large subunit among Eukaryotes and is similar in sequence to DNA ligases. This domain is missing in the large-subunit homolog in M. jannaschii. The remaining domains in the two M. jannaschii genes are well-conserved relative to the eukaryotic homologs. Two

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features of the sequence similarity in these domains are of particular interest. First, domain II (an ATPase domain) of the small-subunit homolog is split between two highly conserved amino acids (lysine and threonine) by an intervening sequence of unknown function. Second, the sequence of domain VI has regions that are useful for distinguishing between bacterial and eukaryotic rfc proteins (Bernard, A., et al., EMBO J. 6:4219 (1987); Cullman, G., et al., Molec. Cell Biol. 15:4661 (1995); Uemori, T., et al., J. Bacteriol. 117:2164 (1995); Delarue, M., et al., Prot. Engineer. 3:461 (1990); Gavin. K.A., et al., Science 270:1667 (1995)); the rfc sequence for M. jannaschii shares the characteristic eukaryotic signature in this domain.

We have attempted to identify an origin of replication by searching the M. jannaschii genome sequence with a variety of bacterial and eukaryotic replication-origin consensus sequences. Searches with oriC, ColE1, and autonomously replicating sequences from yeast (Bernard, A., et al., EMBO J. 6:4219 (1987); Cullman, G., et al., Molec. Cell Biol. 15:4661 (1995); Uemori, T., et al., J. Bacteriol. 117:2164 (1995); Delarue, M., et al., Prot. Engineer. 3:461 (1990); Gavin, K.A., et al., Science 270:1667 (1995)) did not identify an origin of replication. With respect to the related cellular processes of replication initiation and cell division, the M. jannaschii genome contains two genes that are putative homologs of Cdc54, a yeast protein that belongs to a family of putative DNA replication initiation proteins (Whitbred, L.A. and Dalton, S., Gene 155:113 (1995)). A third potential regulator of cell division in M. jannaschii is 55% similar at the amino acid level to pelota, a Drosophila protein involved in the regulation of the early phases of meiotic and mitotic cell division (Eberhart, C.G. and Wasserman, S.A., Development 121:3477 (1995)).

In contrast to the putative rfc complex and the initiation of DNA replication, the cell division proteins from *M. jannaschii* most resemble their bacterial counterparts (Rothfield, L.I. and Zhao, C.R., *Cell 84*:183 (1996); Lutkenhaus, J., *Curr. Opp. Gen. Devel. 3*:783 (1993)). Two genes similar to that encoding FtsZ, a ubiquitous bacterial protein, are found in *M. jannaschii*. FtsZ

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is a polymer-forming, guanosine triphosphate (GTP)-hydrolyzing protein with tubulin-like elements; it is localized to the site f septation and forms a constricting ring between the dividing cells. One gene similar to FtsJ, a bacterial cell division protein of undetermined function, also is found in *M. jannaschii*. Three additional genes (MinC, MinD, and MinE) function in concert in Bacteria to determine the site of septation during cell division. In *M. jannaschii*, three MinD-like genes were identified, but none for MinC or MinE. Neither spindle-associated proteins characteristic of eukaryotic cell division nor bacterial mechanochemical enzymes necessary for partitioning the condensed chromosomes were detected in the *M. jannaschii* genome. Taken together, these observations raise the possibility that cell division in *M. jannaschii* might occur via a mechanism specific for the Archaea.

The structural and functional conservation of the signal peptide of secreted proteins in Archaea, Bacteria, and Eukaryotes suggests that the basic mechanisms of membrane targeting and translocation may be similar among all three domains of life. The secretory machinery of M. jannaschii appears a rudimentary apparatus relative to that of bacterial and eukaryotic systems and consists of (i) a signal peptidase (SP) that cleaves the signal peptide of translocating proteins, (ii) a preprotein translocase that is the major constituent of the membrane-localized translocation channel, (iii) a ribonucleoprotein complex (signal recognition particle, SRP) that binds to the signal peptide and guides nascent proteins to the cell membrane, and (iv) a docking protein that acts as a receptor for the SRP. The 7S RNA component of the SRP from M. jannaschii shows a highly conserved structural domain shared by other Archaea, Bacteria, and Eukaryotes (Kaine, B.P. and Merkel, V.L., J. Bacteriol. 171:4261 (1989); Poritz, M.A. et al., Cell 55:4 (1988)). However, the predicted secondary structure of the 7S RNA SRP component in Archaea is more like that found in Eukaryotes than in Bacteria (Kaine, B.P. and Merkel, V.L., J. Bacteriol. 171:4261 (1989); Poritz, M.A. et al., Cell 55:4 (1988)). The SP and docking proteins from M. jannaschii are most similar to their eukaryotic counterparts; the translocase is most similar to the SecY translocation-associated protein in Escherichia coli.

A second distinct signal peptide is found in the flagellin genes of M. jannaschii. Alignment of flagellin genes from M. voltae (Faguy, D.M., et al., Can. J. Microbiol. 40:67 (1994); Kalmokoff, M.L., et al., Arch. Microbiol. 157:481 (1992)) and M. jannaschii reveals a highly conserved NH₂-terminus (31 of the first 50 residues are identical in all of the mature flagellins). The peptide sequence of the M. jannaschii flagellin indicates that the protein is cleaved after the canonical Gly-12 position, and it is proposed to be similar to type-IV pilins of Bacteria (Faguy, D.M., et al., Can. J. Microbiol. 40:67 (1994); Kalmokoff, M.L., et al., Arch. Microbiol. 157:481 (1992)).

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Five histone genes are present in the *M. jannaschii* genome—three on the main chromosome and two on the large ECE. These genes are homologs of eukaryotic histones (H2a, H2b, H3, and H4) and of the eukaryotic transcription-related CAAT-binding factor CBF-A (Sandman, K., et al., Proc. Natl. Acad. Sci. USA 87:5788 (1990)). The similarity between archaeal and eukaryotic histones suggests that the two groups of organisms resemble one another in the roles histones play both in genome supercoiling dynamics and in gene expression. The five *M. jannaschii* histone genes show greatest similarity among themselves even though a histone sequence is available from the closely related species, *Methanococcus voltae*. This intraspecific similarity suggests that the gene duplications that produced the five histone genes occurred on the *M. jannaschii* lineage per se.

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Self-splicing portions of a peptide sequence that generally encode a DNA endonuclease activity are called inteins, in analogy to introns (Kane, P.M., et al., Science 250:651 (1990); Hirata, R., et al., J. Biol. Chem. 265:6726 (1990); Cooper, A. and Stevens, T., TIBS 20:351 (1995); Xu, M.Q., et al., Cell 75:1371 (1993); Perler et al., Proc. Natl. Acad. Sci. USA 89:5577 (1992); Cooper et al., EMBO J. 12:2575 (1993); Michel et al., Biochimie 64:867 (1992); Pietrokovski S., Prot. Sci. 3:2340 (1994). Most inteins in the M. jannaschii genome were identified by (i) similarity of the bounding exteins to other proteins, (ii) similarity of the inteins to those previously described, (iii) presence of the dodecapeptide endonuclease motifs, and (iv) canonical intein-extein junction sequences. In two

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instances (MJ0832 and MJ0043), the similarity to other database sequences did not unambigu usly define the NH₂-terminal extein-intein junction, so it was necessary to rely on consensus sequences to select the putative site. The inteins in MJ1042 and MJ0542 have previously uncharacterized COOH-terminal splice junctions, GNC and FNC, respectively).

The sequences remaining after an intein is excised are called exteins, in analogy to exons. Exteins are spliced together after the excision of one or more inteins to form functional proteins. The biological significance and role of inteins are not clearly understood (Kane, P.M., et al., Science 250:651 (1990); Hirata, R., et al., J. Biol. Chem. 265:6726 (1990); Cooper, A. and Stevens, T., TIBS 20:351 (1995); Xu, M.Q., et al., Cell 75:1371 (1993); Perler et al., Proc. Natl. Acad. Sci. USA 89:5577 (1992); Cooper et al., EMBO J. 12:2575 (1993); Michel et al., Biochimie 64:867 (1992); Pietrokovski S., Prot. Sci. 3:2340 (1994)). Fourteen genes in the M. jannaschii genome contain 18 putative inteins, a significant increase in the approximately 10 intein-containing genes that have been described (Kane, P.M., et al., Science 250:651 (1990); Hirata, R., et al., J. Biol. Chem. 265:6726 (1990); Cooper, A. and Stevens, T., TIBS 20:351 (1995); Xu, M.Q., et al., Cell 75:1371 (1993); Perier et al., Proc. Natl. Acad. Sci. USA 89:5577 (1992); Cooper et al., EMBO J. 12:2575 (1993); Michel et al., Biochimie 64:867 (1992); Pietrokovski S., Prot. Sci. 3:2340 (1994)) (Table 4). The only previously described inteins in the Archaea are in the DNA polymerase genes of the Thermococcales (Kane, P.M., et al., Science 250:651 (1990); Hirata, R., et al., J. Biol. Chem. 265:6726 (1990); Cooper, A. and Stevens, T., TIBS 20:351 (1995); Xu, M.Q., et al., Cell 75:1371 (1993); Perler et al., Proc. Natl. Acad. Sci. USA 89:5577 (1992); Cooper et al., EMBO J. 12:2575 (1993); Michel et al., Biochimie 64:867 (1992); Pietrokovski S., Prot. Sci. 3:2340 (1994)). The M. jannaschii DNA polymerase gene has two inteins in the same locations as those in Pyrococcus sp. strain KOD1. In this case, the exteins exhibit 46% amino acid identity, whereas intein 2 f the two organisms has only 33% identity. This divergence suggests that intein 2 has not been recently (laterally) transferred between the Thermococcales and M. jannaschii. In contrast, the intein 1

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sequences are 56% identical, more than that of the gene containing them, and comparable to the divergence of inteins within the Thermococcales. This high degree of sequence similarity might be the result of an intein transfer more recent than the splitting of these species. The large number of inteins found in *M. jannaschii* led us to question whether these inteins have been increasing in number by moving within the genome. If this were so, we would expect to find some pairs of inteins that are particularly similar. Comparisons of these and other available intein sequences showed that the closest relationships are those noted above linking the DNA polymerase inteins to correspondingly positioned elements in the Thermococcales. Within *M. jannaschii*, the highest identity observed was 33% for a 380-bp portion of two inteins. This finding suggests that the diversification of the inteins predates the divergence of the *M. jannaschii* and *Pyrococcus* DNA polymerases.

Three families of repeated genetic elements were identified in the M. jannaschii genome. Within two of the families, at least two members were identified as ORFs with a limited degree of sequence similarity to bacterial transposases. Members of the first family, designated ISAMJ1, are repeated 10 times on the main chromosome and once on the large ECE (Fig. 2). There is no sequence similarity between the IS elements in M. jannaschii and the ISMI mobile element described previously for Methanobrevibacter smithii (Hamilton, P.T. et al., Mol. Gen. Genet. 200:47 (1985)). Two members of this family were identified as ORFs and are 27% identical (at the amino acid sequence level) to a transposase from Bacillus thuringiensis (IS240; GenBank accession number M23741). Relative to these two members, the remaining members of the ISAMJI family are missing an internal region of several hundred nucleotides (Fig. 2). With one exception, all members of this family end with 16-bp terminal inverted repeats typical of insertion sequences. One member is missing the terminal repeat at its 5' end. The second family consists of two ORFs that are identical across 928 bp. The ORFs are 23% identical at the amino acid sequence level to the COOH-terminus of a transposase from Lactococcus lactis (IS982; GenBank

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accession number L34754). Neither of the members of the second family contains terminal inverted repeats.

Eighteen copies of the third family of repeated genetic structures (Fig. 3) are distributed fairly evenly around the *M. jannaschii* genome. Unlike the genetic elements described above, none of the components of this repeat unit appears to have coding potential. The repeat structure is composed of a long segment followed by one to 25 tandem repetitions of a short segment. The short segments are separated by sequence that is unique within and among the complete repeat structure. Three similar types of short segments were identified; however, the type of short repeat is consistent within each repeat structure, except for variation of the last short segment in six repeat structures. Similar tandem repeats of short segments have been observed in Bacteria and other Archaea (Mojica, F.J.M., et al., Mol. Micro. 17:85 (1995)) and have been hypothesized to participate in chromosome partitioning during cell division.

The 16-kbp ECE from *M. jannaschii* contains 12 ORFs, none of which had a significant full-length match to any published sequence. The 58-kbp ECE contains 44 predicted protein-coding regions, 5 of which had matches to genes in the database. Two of the genes are putative archaeal histones, one is a sporulation-related protein (SOJ protein), and two are type I restriction modification enzymes. There are several instances in which predicted protein-coding regions or repeated genetic elements on the large ECE have similar counterparts on the main chromosome of *M. jannaschii*. The degree of nucleotide sequence similarity between genes present on both the ECE and the main chromosome ranges from 70 to 90%, suggesting that there has been relatively recent exchange of at least some genetic material between the large ECE and the main chromosome.

All the predicted protein-coding regions from *M. jannaschii* were searched against each other in order to identify families of paralogous genes (genes related by gene duplication, not speciation). The initial criterion for grouping paralogs was >30% amino acid sequence identity over 50 consecutive amino acid residues. Groups of predicted protein-coding regions were then

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aligned and inspected individually to ensure that the sequence similarity extended over most of their lengths. This curatorial process resulted in the identification of more than 100 gene families, half of which have no database matches. The largest identified gene family (16 members: MJ0625, MJECL28, MJ1076, MJ1006, MJ1659, MJ0075, MJ1609, MJECL19, MJECL18, MJ0147, MJ0801, MJ1301, MJ0632, MJ1010, MJ0074, and MJ0439) contains almost 1% of the total predicted protein-coding regions in *M. jannaschii*.

Despite the availability for comparison of two complete bacterial genomes and several hundred megabase pairs of eukaryotic sequence data, the majority of genes in M. jannaschii cannot be identified on the basis of sequence similarity. Previous evidence for the shared common ancestry of the Archaeal and Eukaryotic was based on a small set gene sequences (Iwabe, N., et al., Proc. Natl. Acad. Sci. USA 86:9355 (1989); Gogarten J.P., et al., Proc. Natl. Acad. Sci. USA 86:6661 (1989); Brown, J.R. and Doolittle, W.F., Proc. Natl. Acad. Sci. USA 92:2441 (1995)). The complete genome of M. jannaschii allows us to move beyond a "gene by gene" approach to one that encompasses the larger picture of metabolic capacity and cellular systems. The anabolic genes of M. jannaschii (especially those related to energy production and nitrogen fixation) reveal an ancient metabolic world shared largely by Bacteria and Archaea. That many basic autotrophic pathways appear to have a common evolutionary origin suggests that the most recent universal common ancestor to all three domains of extant life had the capacity for autotrophy. The Archaea and Bacteria also share structural and organizational features that the most recent universal prokaryotic ancestors also likely possessed, such as circular genomes and genes organized as operons. In contrast, the cellular information-processing and secretion systems in M. jannaschii demonstrate the common ancestry of Eukaryotes and Archaea. Although there are components of these systems are present in all three domains, apparent refinement over time-especially transcription translation-indicate that the Archaea and Eukaryotes share a common evolutionary trajectory independent of the lineage of Bacteria.

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Example 2

Preparation of PCR Primers and Amplification of DNA

Various fragments of the *Methanococcus jannaschii* genome, such as those disclosed in Tables 2(a), 2(b) and 3 can be used, in accordance with the present invention, to prepare PCR primers. The PCR primers are preferably at least 15 bases, and more preferably at least 18 bases in length. When selecting a primer sequence, it is preferred that the primer pairs have approximately the same G/C ratio, so that melting temperatures are approximately the same. The PCR primers are useful during PCR cloning of the ORFs described herein.

Example 3

Gene expression from DNA Sequences Corresponding to ORFs

A fragment of the *Methanococcus jannaschii* genome (preferably, a protein-encoding sequence) provided in Tables 2(a), 2(b) or 3 is introduced into an expression vector using conventional technology (techniques to transfer cloned sequences into expression vectors that direct protein translation in mammalian, yeast, insect or bacterial expression systems are well known in the art). Commercially available vectors and expression systems are available from a variety of suppliers including Stratagene (La Jolla, California), Promega (Madison, Wisconsin), and Invitrogen (San Diego, California). If desired, to enhance expression and facilitate proper protein folding, the codon context and codon pairing of the sequence may be optimized for the particular expression organism, as explained by Hatfield *et al.*, U.S. Pat. No. 5,082,767, which is hereby incorporated by reference.

The following is provided as one exemplary method to generate polypeptide(s) from a cloned ORF of the *Methanococcus* gen me whose sequence is provided in SEQ ID NOS: 1, 2 and 3. A poly A sequence can be

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added to the construct by, for example, splicing ut the poly A sequence from pSG5 (Stratagene) using BgII and SaII restriction endonuclease enzymes and incorporating it into the mammalian expression vector pXT1 (Stratagene) for use in eukaryotic expression systems. pXT1 contains the LTRs and a portion of the gag gene from Moloney Murine Leukemia Virus. The position of the LTRs in the construct allow efficient stable transfection. The vector includes the Herpes Simplex thymidine kinase promoter and the selectable neomycin gene. The Methanococcus DNA is obtained by PCR from the bacterial vector using oligonucleotide primers complementary to the Methanococcus DNA and containing restriction endonuclease sequences for PstI incorporated into the 5' primer and Bg/II at the 5' end of the corresponding Methanococcus DNA 3' primer, taking care to ensure that the Methanococcus DNA is positioned such that its followed with the poly A sequence. The purified fragment obtained from the resulting PCR reaction is digested with PstI, blunt ended with an exonuclease, digested with Bg/II, purified and ligated to pXT1, now containing a poly A sequence and digested BgIII.

The ligated product is transfected into mouse NIH 3T3 cells using Lipofectin (Life Technologies, Inc., Grand Island, New York) under conditions outlined in the product specification. Positive transfectants are selected after growing the transfected cells in 600 ug/ml G418 (Sigma, St. Louis, Missouri). The protein is preferably released into the supernatant. However if the protein has membrane binding domains, the protein may additionally be retained within the cell or expression may be restricted to the cell surface.

Since it may be necessary to purify and locate the transfected product, synthetic 15-mer peptides synthesized from the predicted *Methanococcus* DNA sequence are injected into mice to generate antibody to the polypeptide encoded by the *Methanococcus* DNA.

If antibody production is not possible, the *Methanococcus* DNA sequence is additionally incorporated into eukaryotic expression vectors and expressed as a chimeric with, for example, \(\beta \)-globin. Antibody to \(\beta \)-globin is used to purify the chimeric. Corresponding protease cleavage sites engineered between the \(\beta \)-globin

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gene and the *Methanococcus* DNA are then used to separate the two polypeptide fragments from ne another after translation. One useful expression vector for generating β-globin chimerics is pSG5 (Stratagene). This vector encodes rabbit β-globin. Intron II of the rabbit β-globin gene facilitates splicing of the expressed transcript, and the polyadenylation signal incorporated into the construct increases the level of expression. These techniques as described are well known to those skilled in the art of molecular biology. Standard methods are available from the technical assistance representatives from Stratagene, Life Technologies, Inc., or Promega. Polypeptides may additionally be produced from either construct using in vitro translation systems such as In vitro ExpressTM Translation Kit (Stratagene).

Example 4

E. coli Expression of a M. jannaschii ORF and protein purification

A M. jannaschii ORF described in Table 2(a), 2(b), or 3 is selected and amplified using PCR oligonucleotide primers designed from the nucleotide sequences flanking the selected ORF and/or from portions of the ORF's NH₂- or COOH-terminus. Additional nucleotides containing restriction sites to facilitate cloning are added to the 5' and 3' sequences, respectively.

The restriction sites are selected to be convenient to restriction sites in the bacterial expression vector pD10 (pQE9), which is used for bacterial expression. (Qiagen, Inc. 9259 Eton Avenue, Chatsworth, CA, 91311). [pD10]pQE9 encodes ampicillin antibiotic resistance ("Ampr") and contains a bacterial origin of replication ("ori"), an IPTG inducible promoter, a ribosome binding site ("RBS"), a 6-His tag and restriction enzyme sites.

The amplified *M. jannaschii* DNA and the vector pQE9 both are digested with Sall and Xbal and the digested DNAs are then ligated together. Insertion of the *M. jannaschii* DNA into the restricted pQE9 vector places the *M. jannaschii* coding region downstream of and operably linked to the vector's IPTG-inducible

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promoter and in-frame with an initiating AUG appropriately positioned for translation of the *M. jannaschii* protein.

The ligation mixture is transformed into competent *E. coli* cells using standard procedures. Such procedures are described in Sambrook *et al.*, Molecular Cloning: a Laboratory Manual, 2nd Ed.; Cold Spring Harbor Laboratory Press, Cold Spring Harbor, N.Y. (1989). *E. coli* strain M15/rep4, containing multiple copies of the plasmid pREP4, which expresses lac repressor and confers kanamycin resistance ("Kan"), is used in carrying out the illustrative example described herein. This strain, which is only one of many that are suitable for expressing *M. jannaschii* protein, is available commercially from Qiagen.

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Transformants are identified by their ability to grow on LB plates in the presence of ampicillin and kanamycin. Plasmid DNA is isolated from resistant colonies and the identity of the cloned DNA confirmed by restriction analysis. Clones containing the desired constructs are grown overnight ("O/N") in liquid culture in LB media supplemented with both ampicillin (100 μ g/ml) and kanamycin (25 μ g/ml).

The O/N culture is used to inoculate a large culture, at a dilution of approximately 1:100 to 1:250. The cells are grown to an optical density at 600nm ("OD600") of between 0.4 and 0.6. Isopropyl-B-D-thiogalactopyranoside ("IPTG") is then added to a final concentration of 1 mM to induce transcription from *lac* repressor sensitive promoters, by inactivating the *lac*I repressor. Cells subsequently are incubated further for 3 to 4 hours. Cells then are harvested by centrifugation and disrupted, by standard methods. Inclusion bodies are purified from the disrupted cells using routine collection techniques, and protein is solubilized from the inclusion bodies into 8M urea. The 8M urea solution containing the solubilized protein is passed over a PD-10 column in 2X phosphate-buffered saline ("PBS"), thereby removing the urea, exchanging the buffer and refolding the protein. The protein is purified by a further step of chromatography to remove endotoxin followed by sterile filtration. The sterile filtered protein preparation is stored in 2X PBS at a concentration of 95 µ/ml.

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Example 5

Cloning and Expression of a M. jannaschii protein in a Baculovirus Expression System

A M. jannaschii ORF described in Table 2(a), 2(b), or 3 is selected and amplified as above. The amplified DNA is isolated from a 1% agarose gel using a commercially available kit ("Geneclean," BIO 101 Inc., La Jolla, Ca.). The DNA then is digested with XbaI and again purified on a 1% agarose gel. This DNA is designated herein as F2.

The vector pA2-GP is used to express the *M. jannaschii* protein in the baculovirus expression system as described in Summers *et al.*, A Manual of Methods for Baculovirus Vectors and Insect Cell Culture Procedures. Texas Agricultural Experimental Station Bulletin No. 1555 (1987). The pA2-GP expression vector contains the strong polyhedrin promoter of the *Autographa californica* nuclear polyhedrosis virus (AcMNPV) followed by convenient restriction sites. The signal peptide of AcMNPV gp67, including the N-terminal methionine, is located just upstream of a BamHI site. The polyadenylation site from the simian virus 40 ("SV40") is used for efficient polyadenylation. For an easy selection of recombinant virus, the beta-galactosidase gene from *E. coli* is inserted in the same orientation as the polyhedrin promoter and is followed by the polyadenylation signal of the polyhedrin genc. The polyhedrin sequences are flanked at both sides by viral sequences for cell-mediated homologous recombination with wild-type viral DNA to generate viable virus that express the cloned polynucleotide.

Many other baculovirus vectors could be used in place of pA2-GP, such as pAc373, pVL941 and pAcIM1 provided, as those of skill readily will appreciate, that construction provides appropriately located signals for transcription, translation, trafficking and the like, such as an in-frame AUG and a signal peptide, as required. Such vectors are described in Luckow et al., Virology 170: 31-39, among others.

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The plasmid is digested with the restriction enzyme Xbal and then is dephosphorylated using calf intestinal phosphatase, using routine procedures known in the art. The DNA is then isolated from a 1% agarose gel using a commercially available kit ("Geneclean" BIO 101 Inc., La Jolla, Ca.). This vector DNA is designated herein "V".

Fragment F2 and the dephosphorylated plasmid V2 are ligated together with T4 DNA ligase. E. coli HB101 cells are transformed with ligation mix and spread on culture plates. Bacteria are identified that contain the plasmid with the M. jannaschii gene by digesting DNA from individual colonies using Xbal and then analyzing the digestion product by gel electrophoresis. The sequence of the cloned fragment is confirmed by DNA sequencing. This plasmid is designated herein pBacM. jannaschii.

5 μg of the plasmid pBac*M. jannaschii* is co-transfected with 1.0 μg of a commercially available linearized baculovirus DNA ("BaculoGold™ baculovirus DNA", Pharmingen, San Diego, CA.), using the lipofection method described by Felgner *et al.*, Proc. Natl. Acad. Sci. USA 84: 7413-7417 (1987). 1μg of BaculoGold™ virus DNA and 5 μg of the plasmid pBac*M. jannaschii* are mixed in a sterile well of a microtiter plate containing 50 μl of serum-free Grace's medium (Life Technologies Inc., Gaithersburg, MD). Afterwards 10 μl Lipofectin plus 90 μl Grace's medium are added, mixed and incubated for 15 minutes at room temperature. Then the transfection mixture is added drop-wise to Sf9 insect cells (ATCC CRL 1711) seeded in a 35 mm tissue culture plate with 1 ml Grace's medium without serum. The plate is rocked back and forth to mix the newly added solution. The plate is then incubated for 5 hours at 27°C. After 5 hours the transfection solution is removed from the plate and 1 ml of Grace's insect medium supplemented with 10% fetal calf serum is added. The plate is put back into an incubator and cultivation is continued at 27°C for four days.

After four days the supernatant is collected and a plaque assay is performed, as described by Summers and Smith, cited above. An agarose gel with "Blue Gal" (Life Technologies Inc., Gaithersburg) is used to allow easy identification and isolation of gal-expressing clones, which produce blue-stained

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plaques. (A detailed description f a "plaque assay" of this type can also be found in the user's guide for insect cell culture and baculovirology distributed by Life Technologies Inc., Gaithersburg, page 9-10).

Four days after serial dilution, the virus is added to the cells. After appropriate incubation, blue stained plaques are picked with the tip of an Eppendorf pipette. The agar containing the recombinant viruses is then resuspended in an Eppendorf tube containing 200 µl of Grace's medium. The agar is removed by a brief centrifugation and the supernatant containing the recombinant baculovirus is used to infect Sf9 cells seeded in 35 mm dishes. Four days later the supernatants of these culture dishes are harvested and then they are stored at 4°C. A clone containing properly inserted hESSB I, II and III is identified by DNA analysis including restriction mapping and sequencing. This is designated herein as V-M. jannaschii.

Sf9 cells are grown in Grace's medium supplemented with 10% heat-inactivated FBS. The cells are infected with the recombinant baculovirus V-M. jannaschii at a multiplicity of infection ("MOI") of about 2 (about 1 to about 3). Six hours later the medium is removed and is replaced with SF900 II medium minus methionine and cysteine (available from Life Technologies Inc., Gaithersburg). 42 hours later, 5 µCi of ³⁵S-methionine and 5 µCi ³⁵S-cysteine (available from Amersham) are added. The cells are further incubated for 16 hours and then they are harvested by centrifugation, lysed and the labeled proteins are visualized by SDS-PAGE and autoradiography.

Example 6

Cloning and Expression in Mammalian Cells

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Most of the vectors used for the transient expression of a *M. jannaschii* gene in mammalian cells should carry the SV40 origin of replication. This allows the replication of the vector to high copy numbers in cells (e.g., COS cells) which

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express the T antigen required f r the initiation of viral DNA synthesis. Any other mammalian cell line can also be utilized for this purpose.

A typical mammalian expression vector contains the promoter element, which mediates the initiation of transcription of mRNA, the protein-coding sequence, and signals required for the termination of trancription and polyadenylation of the transcript. Additional elements include enhancers, Kozak sequences and intervening sequences flanked by donor and acceptor sites for RNA splicing. Highly efficient transcription can be achieved with the early and late promoters from SV40, the long terminal repeats (LTRs) from Retroviruses, e.g., RSV, HTLVI, HIVI and the early promoter of the cytomegalovirus (CMV). However, cellular signals can also be used (e.g., human actin promoter). Suitable expression vectors for use in practicing the present invention include, for example, vectors such as pSVL and pMSG (Pharmacia, Uppsala, Sweden), pRSVcat (ATCC 37152), pSV2dhfr (ATCC 37146) and pBC12MI (ATCC 67109). Mammalian host cells that could be used include, human Hela, 283, H9 and Jurkart cells, mouse NIH3T3 and C127 cells, Cos 1, Cos 7 and CV1, African green monkey cells, quail QC1-3 cells, mouse L cells and Chinese hamster ovary cells.

Alternatively, the gene can be expressed in stable cell lines that contain the gene integrated into a chromosome. The co-transfection with a selectable marker such as dhfr, gpt, neomycin, hygromycin allows the identification and isolation of the transfected cells.

The transfected gene can also be amplified to express large amounts of the encoded protein. The DHFR (dihydrofolate reductase) is a useful marker to develop cell lines that carry several hundred or even several thousand copies of the gene of interest. Another useful selection marker is the enzyme glutamine synthase (GS) (Murphy et al., Biochem J. 227:277-279 (1991); Bebbington et al., Bio/Technology 10:169-175 (1992)). Using these markers, the mammalian cells are grown in selective medium and the cells with the highest resistance are selected. These cell lines contain the amplified gene(s) integrated into a

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chromosome. Chinese hamster ovary (CHO) cells are often used for the production of proteins.

The expression vectors pC1 and pC4 contain the strong promoter (LTR) of the Rous Sarcoma Virus (Cullen et al., Molecular and Cellular Biology, 438-447 (March, 1985)) plus a fragment of the CMV-enhancer (Boshart et al., Cell 41:521-530 (1985)). Multiple cloning sites, e.g., with the restriction enzyme cleavage sites BamHI, XbaI and Asp718, facilitate the cloning of the gene of interest. The vectors contain in addition the 3' intron, the polyadenylation and termination signal of the rat preproinsulin gene.

Example 6(a): Cloning and Expression in COS Cells

The expression plasmid, p*M jannaschii* HA, is made by cloning a cDNA encoding a *M. jannaschii* protein into the expression vector pcDNAI/Amp (which can be obtained from Invitrogen, Inc.).

The expression vector pcDNAI/amp contains: (1) an E. coli origin of replication effective for propagation in E. coli and other prokaryotic cells; (2) an ampicillin resistance gene for selection of plasmid-containing prokaryotic cells; (3) an SV40 origin of replication for propagation in eukaryotic cells; (4) a CMV promoter, a polylinker, an SV40 intron, and a polyadenylation signal arranged so that a cDNA conveniently can be placed under expression control of the CMV promoter and operably linked to the SV40 intron and the polyadenylation signal by means of restriction sites in the polylinker.

A DNA fragment encoding the *M. jannaschii* protein and an HA tag fused in frame to its 3' end is cloned into the polylinker region of the vector so that recombinant protein expression is directed by the CMV promoter. The HA tag corresponds to an epitope derived from the influenza hemagglutinin protein described by Wilson *et al.*, *Cell 37:767* (1984). The fusion of the HA tag to the target protein allows easy detection of the recombinant protein with an antibody that recognizes the HA epitope.

The PCR amplified DNA fragment (generated as described above) and the vector, pcDNAI/Amp, are digested with HindIII and XhoI and then ligated. The ligation mixture is transformed into *E. coli* strain SURE (available from Stratagene Cloning Systems, 11099 North Torrey Pines Road, La Jolla, CA 92037), and the transformed culture is plated on ampicillin media plates which then are incubated to allow growth of ampicillin resistant colonies. Plasmid DNA is isolated from resistant colonies and examined by restriction analysis and gel sizing for the presence of the *M. jannaschii* protein-encoding fragment.

For expression of recombinant *M. jannaschii*, COS cells are transfected with an expression vector, as described above, using DEAE-DEXTRAN, as described, for instance, in Sambrook *et al.*, Molecular Cloning: a Laboratory Manual, Cold Spring Laboratory Press, Cold Spring Harbor, New York (1989). Cells are incubated under conditions for expression of *M. jannaschii* protein by the vector.

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Expression of the *M. jannaschii* HA fusion protein is detected by radiolabelling and immunoprecipitation, using methods described in, for example Harlow *et al.*, Antibodies: A Laboratory Manual, 2nd Ed.; Cold Spring Harbor Laboratory Press, Cold Spring Harbor, New York (1988). To this end, two days after transfection, the cells are labeled by incubation in media containing ³⁵S-cysteine for 8 hours. The cells and the media are collected, and the cells are washed and the lysed with detergent-containing RIPA buffer: 150 mM NaCl, 1% NP-40, 0.1% SDS, 1% NP-40, 0.5% DOC, 50 mM TRIS, pH 7.5, as described by Wilson *et al.* cited above. Proteins are precipitated from the cell lysate and from the culture media using an HA-specific monoclonal antibody. The precipitated proteins then are analyzed by SDS-PAGE gels and autoradiography. An expression product of the expected size is seen in the cell lysate, which is not seen in negative controls.

Example 6(b): Cloning and Expression in CHO Cells

The vector pC1 is used for the expression of a M. jannaschii protein. Plasmid pC1 is a derivative of the plasmid pSV2-dhfr [ATCC Accession No. 37146]. Both plasmids contain the mouse DHFR gene under control of the SV40 early promoter. Chinese hamster ovary- or other cells lacking dihydrofolate activity that are transfected with these plasmids can be selected by growing the cells in a selective medium (alpha minus MEM, Life Technologies) supplemented with the chemotherapeutic agent methotrexate. The amplification of the DHFR genes in cells resistant to methotrexate (MTX) has been well documented (see. e.g., Alt, F.W., Kellems, R.M., Bertino, J.R., and Schimke, R.T., 1978, J. Biol. Chem. 253:1357-1370, Hamlin, J.L. and Ma, C. 1990, Biochem. et Biophys. Acta, 1097:107-143, Page, M.J. and Sydenham, M.A. 1991, Biotechnology Vol. 9:64-68). Cells grown in increasing concentrations of MTX develop resistance to the drug by overproducing the target enzyme, DHFR, as a result of amplification of the DHFR gene. If a second gene is linked to the DHFR gene it is usually co-amplified and over-expressed. It is state of the art to develop cell lines carrying more than 1,000 copies of the genes. Subsequently, when the methotrexate is withdrawn, cell lines contain the amplified gene integrated into the chromosome(s).

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Plasmid pC1 contains for the expression of the gene of interest a strong promoter of the long terminal repeat (LTR) of the Rouse Sarcoma Virus (Cullen, et al., Molecular and Cellular Biology, March 1985:438-4470) plus a fragment isolated from the enhancer of the immediate early gene of human cytomegalovirus (CMV) (Boshart et al., Cell 41:521-530, 1985). Downstream of the promoter are the following single restriction enzyme cleavage sites that allow the integration of the genes: BamHI, Pvull, and Nrul. Behind these cloning sites the plasmid contains translational stop codons in all three reading frames followed by the 3' intron and the polyadenylation site of the rat preproinsulin gene. Other high efficient promoters can also be used for the expression, e.g., the human β-actin promoter, the SV40 early or late promoters or the long terminal

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repeats from other retroviruses, e.g., HIV and HTLVI. F r the polyadenylation of the mRNA other signals, e.g., from the human growth hormone or globin genes can be used as well.

Stable cell lines carrying the gene of interest integrated into the chromosomes can also be selected upon co-transfection with a selectable marker such as gpt, G418 or hygromycin. It is advantageous to use more than one selectable marker in the beginning, e.g., G418 plus methotrexate.

The plasmid pC1 is digested with the restriction enzyme BamHl and then dephosphorylated using calf intestinal phosphates by procedures known in the art. The vector is then isolated from a 1% agarose gel.

The *M. jannaschii* protein-encoding sequence is is amplified using PCR oligonucleotide primers as described above. An efficient signal for initiation of translation in eukaryotic cells, as described by Kozak, M., J. Mol. Biol. 196:947-950 (1987) is appropriately located in the vector portion of the construct. The amplified fragments are isolated from a 1% agarose gel as described above and then digested with the endonucleases BamHI and Asp718 and then purified again on a 1% agarose gel.

The isolated fragment and the dephosphorylated vector are then ligated with T4 DNA ligase. *E. coli* HB101 cells are then transformed and bacteria identified that contained the plasmid pC1 inserted in the correct orientation using the restriction enzyme BamHI. The sequence of the inserted gene is confirmed by DNA sequencing.

Transfection of CHO-DHFR-cells

Chinese hamster ovary cells lacking an active DHFR enzyme are used for transfection. 5 µg of the expression plasmid C1 are cotransfected with 0.5 µg of the plasmid pSVneo using the lipofecting method (Felgner et al., supra). The plasmid pSV2-neo contains a dominant selectable marker, the gene neo from Tn5 encoding an enzyme that confers resistance to a group of antibiotics including G418. The cells are seeded in alpha minus MEM supplemented with 1 mg/ml

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G418. After 2 days, the cells are trypsinized and seeded in hybridoma cloning plates (Greiner, Germany) and cultivated from 10-14 days. After this period, single clones are trypsinized and then seeded in 6-well petri dishes using different concentrations of methotrexate (25 nM, 50 nM, 100 nM, 200 nM, 400 nM). Clones growing at the highest concentrations of methotrexate are then transferred to new 6-well plates containing even higher concentrations of methotrexate (500 nM, 1 μ M, 2 μ M, 5 μ M). The same procedure is repeated until clones grow at a concentration of 100 μ M.

The expression of the desired gene product is analyzed by Western blot analysis and SDS-PAGE.

Example 7

Production of an Antibody to a Methanococcus jannaschii Protein

Substantially pure *M. jannaschii* protein or polypeptide is isolated from the transfected or transformed cells described above using an art-known method. The protein can also be chemically synthesized. Concentration of protein in the final preparation is adjusted, for example, by concentration on an Amicon filter device, to the level of a few micrograms/ml. Monoclonal or polyclonal antibody to the protein can then be prepared as follows:

Monoclonal Antibody Production by Hybridoma Fusion

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Monoclonal antibody to epitopes of any of the peptides identified and isolated as described can be prepared from murine hybridomas according to the classical method of Kohler, G. and Milstein, C., *Nature 256*:495 (1975) or modifications of the methods thereof. Briefly, a mouse is repetitively inoculated with a few micrograms of the selected protein over a period of a few weeks. The mouse is then sacrificed, and the antibody producing cells of the spleen isolated. The spleen cells are fused by means of polyethylene glycol with mouse myeloma

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cells, and the excess unfused cells destroyed by growth of the system in selective media comprising aminopterin (HAT media). The successfully fused cells are diluted and aliquots of the dilution placed in wells of a microtiter plate where growth of the culture is continued. Antibody-producing clones are identified by detection of antibody in the supernatant fluid of the wells by immunoassay procedures, such as ELISA, as originally described by Engvall, E., Meth. Enzymol. 70:419 (1980), and modified methods thereof. Selected positive clones can be expanded and their monoclonal antibody product harvested for use. Detailed procedures for monoclonal antibody production are described in Davis. L. et al. Basic Methods in Molecular Biology Elsevier, New York. Section 21-2 (1989).

Polyclonal Antibody Production by Immunization

Polyclonal antiserum containing antibodies to heterogenous epitopes of a single protein can be prepared by immunizing suitable animals with the expressed protein described above, which can be unmodified or modified to enhance immunogenicity. Effective polyclonal antibody production is affected by many factors related both to the antigen and the host species. For example, small molecules tend to be less immunogenic than other molecules and may require the use of carriers and adjuvant. Also, host animals vary in response to site of inoculations and dose, with both inadequate or excessive doses of antigen resulting in low titer antisera. Small doses (ng level) of antigen administered at multiple intradermal sites appears to be most reliable. An effective immunization protocol for rabbits can be found in Vaitukaitis, J. et al., J. Clin. Endocrinol. Metab. 33:988-991 (1971).

Booster injections can be given at regular intervals, and antiserum harvested when antibody titer thereof, as determined semi-quantitatively, for example, by double immunodiffusion in agar against known concentrations of the antigen, begins to fall (See Ouchterlony, O. et al., Chap. 19 in: Handbook of Experimental Immunology, Wier, D., ed, Blackwell (1973)). Plateau

concentration of antibody is usually in the range of 0.1 to 0.2 mg/ml of serum (about 12 pM). Affinity of the antisera for the antigen is determined by preparing competitive binding curves, as described, for example, by Fisher, D., Chap. 42 in: Manual of Clinical Immunology, second edition, Rose and Friedman, (eds.), Amer. Soc. For Microbio., Washington, D.C. (1980).

Antibody preparations prepared according to either protocol are useful in quantitative immunoassays which determine concentrations of antigen-bearing substances in biological samples; they are also used semi-quantitatively or qualitatively to identify the presence of antigen in a biological sample.

Table 2A

			A				
			Amino acid biosynthesis				_
Aromatic amino acid family	ino acid fam	ily					Т
MJ1454	47830	48200					
	200	40390	3-dehydroquinate dehydratase (Escherichia coli)	300	67.9	3	7
MJ0502	1029204	1027915	5-enolpyruvylshikimate 1-nhoenhote armst.	3,5:0	24.0	100	7
MJ1075	456842	458159	the syndia's syndia's (Haemophilus influenzae)	38.2	0.09	1290	
		20120	animanilate synthase, subunit I (Clostridium thermocellum)	52.7	72.1	1317	_
MJ0234	1247181	1246243	anthranilate synthage surhing II' (The			131/	_
MJ0238	1242410	1241016	included maritima)	44.1	64.3	939	
		01/21/21	anthranilate synthase, subunit II" (Thermotoga maritima)	203	3.5		т
MJ0246	1238364	1238660	chorismate mutase surhunit A (Emiliais L. L.	32.0	0.0	495	
MJ0612	070701	20000	Comment (Ciwillia Herbicola)	37.4	59.4	297	
	16/676	57/876	chorismate mutase subunit B (Escherichia coli)				_
MJ1175	357469	358572	Chorismate sumbace (C.	33.2	56.2	1059	-4
0.0017			constructed synechocystis sp)	48	3 99	1104	.7
MJ0918	621924	622682	indole-3-glycerol phosphate synthese (11st-base)			507-	_
MJ0451	1068501	1023046	Similary (Trainoracienium Voicanii)	42.7	67.7	759	
	100001	100/043	N-phosphoribosyl anthranilate isomerase (Haloferax volcanii)	9 17	3		_
MJ0637	904569	905264	prephenate dehydratase // actonomic 1	۲.۱۰۶	67.5	657	
M11084	440622		- Caciococcus lacits	39.3	61.7	969	
100100	449333	448/57	shikimate 5-dehydrogenase (Escherichia coli)				_
MJ1038	502619	501777	Trategraphy	38.9	57.4	77,	
2411017			"This print synthase, subunit alpha (Methanobacterium thermoautotrophicum)	49.8	69.3	843	,
1601037	503929	502808	tryptophan synthase, subunit beta (Acinetobacter calcoaceticus)	T			
				600	707		_

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Aspartate family	lly					
MJ1116	414120	415679	asparagine synthetase {Escherichia coli}	34.0	54.3	1560
MJ1056	476613	476170	asparagine synthetase {Bacillus subtilis}	33.0	54.6	444
MJ1391	132691	133833	aspartate aminotransferase {Sulfolobus solfataricus}	31.0	52.2	1143
MJ0684	859565	860632	aspartate aminotransferase (Sulfolobus solfataricus)	37.8	63.7	1068
MJ0001	1469369	1470142	aspartate aminotransferase (Sulfolobus solfataricus)	39.2	63.8	774
MJ0205	1273947	1274951	aspartate-semialdehyde dehydrogenase {Leptospira interrogans}	50.4	67.2	1005
MJ0571	963902	962544	aspartokinase I (Serratia marcescens)	37.0	56.7	1359
MJ1473	26812	27558	cobalamin-independent methionine synthase {Methanobacterium thermoautotrophicum}	47.7	65.3	747
MJ1097	433957	435159	diaminopimelate decarboxylase (Haemophilus influenzae)	43.2	9.99	1203
MJ1119	412913	412029	diaminopimelate epimerase (Haemophilus influenzae)	36.2	9.99	885
MJ0422	1090629	1091441	dihydrodipicolinate reductase {Haemophilus influenzae}	45.0	64.4	813
MJ0244	1239093	1239776	dihydrodipicolinate synthase (Haemophilus influenzae)	46.6	64.4	684
MJ1003	540278	539106	homoaconitase {Saccharomyces cerevisiae}	35.7	56.9	1173
MJ1602	1563296	1562289	homoserine dehydrogenase {Bacillus subtilis}	40.4	63.2	1008
MJ1104	427241	428128	homoserine kinase (Haemophilus influenzae)	30.1	53.9	80 80 80
MJ0020	1450056	1451210	L-asparaginase I (Haemophilus influenzae)	34.8	53.1	1155

MJ0457	1064285	1063176	Chorine, Line			
MJ1465	16087	20167	occury i-usaiminopimelate desuccinylase (Haemophilus influenzae)	27.0	45.8	1110
	20205	3013/	threonine synthase (Bacillus subtilis)	613	1 - 1	761.
Glutamate family	ımily					0/1
MJ0069	1406333	1405455				
M10791	767716			44.4	65.7	879
	(16/6/	/28637	argininosuccinate lyase (Campylobacter ieiuni)			
MJ0429	1087105	1086023	argininosuccinate synthage (Methanogram)	41.3	65.6	1323
MJ0186	1287178	1288140	olutamate Negetities of	70.2	86.8	1083
MJ1351	172535	174007	of intermedia constitution (Constitution of intermedia constitution of inte	47.4	63.1	963
MJ1346	179417	178068	of the synthesis (NADPH), subunit alpha (Escherichia coli)	40.5	54.0	1473
MJ1096	435486	416500	Successions synthetiase (Methanococcus voltae)	70.5	84.7	1350
		430300	N-acetyl-gamma-glutamyl-phosphate reductase (Bacillus subtilis)	40.4	717	1631
MJ0/21	817148	816045	N-acetylomithine aminotransferase (Anabases)		9:50	1023
MJ0881	664952	665845	omithine carbamovirence of the carbamovirenc	46.7	67.0	69. 5011
Pyruvate family	4		(nalobacterium halobium)	43.0	9.69	894
MJ0503	1027812	1026610	2 iconsonut			
MJ1392	131826	130633	2 :	44.4	61.1	1203
M11271	366614		2-130propyimalate synthase {Anabaena sp.}	43.0	63.1	1194
	410007	726216	3-isopropylmalate dehydratase (Salmonella typhimurium)	1	十	
MJ1277	249421	249807	3-isopropylmalate dehydratase (Clostridium pactameio	十	0.2.0	399
MJ0663	884580	883129	acetolactate conthaca farma culturation in the contract of the	49.5	70.2	387
MJ0277	1207735	1200507	Commercial Se suculit (Forpnyra umbilicalis)	34.5	54.6	1452
		1000000	accionaciare synthase, large subunit (Bacillus subtilis)	603	40.7	1333

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MJ0161	1307199	1307702	acetolactate synthase, small subunit (Bacillus subtilis)	49.4	74.1	504
MJ1008	533323	534132	branched-chain amino acid aminotransferase (Escherichia coli)	42.6	59.0	810
MJ1276	250052	251710	dihydroxy-acid dehydratase {Lactococcus lactis}	44.6	65.1	1659
MJ1195	333450	335003	isopropylmalate synthase (Hacmophilus influenzae)	42.9	63.7	1554
MJ1543	1615932	1614931	ketol-acid reductoisomerase (Bacillus subtilis)	53.7	77.0	1002
Serine family						
MJ1597	1568671	1567445	glycine hydroxymethyltransferase (Methanobacterium thermoautotrophicum)	8.69	80.7	1227
MJ1018	523454	524806	phosphoglycerate dehydrogenase (Bacillus subtilis)	42.7	65.4	1353
MJ1594	1571545	1571039	phosphoserine phosphatase (Haemophilus influenzae)	40.4	62.7	507
MJ0959	580672	581778	serine aminotransferase (Methanobacterium thermoformicicum)	54.5	74.9	1107
Histidine family	ly					
MJ1204	324063	324878	ATP phosphoribosyltransferase {Escherichia coli}	34.0	57.3	816
MJ1456	46532	45354	histidinol dehydrogenase {Lactococcus lactis}	47.6	67.5	1179
MJ0955	586179	585073	histidinol-phosphate aminotransferase (Bacillus subtilis)	37.7	8.09	1107
MJ0698	848921	848364	imidazoleglycerol-phosphate dehydrogenase {Methanobacterium thermoautotrophicum}	51.7	71.2	558
MJ0506	1024803	1025237	imidazoleglycerol-phosphate synthase (amidotransferase) {Lactococcus lactis}	45.6	62.1	435
MJ0411	1101451	1100636	imidazoleglycerol-phosphate synthase (cyclase) {Azospirillum brasilense}	61.5	78.8	816
MJ1430	71328	71047	phosphoribosyl-AMP cyclohydrolase (Methanococcus vannielii)	70.0	86.3	282

MJ0302	1186990	1187208	phosphoribosyl-ATP pyrophosphohydrolase (Azotohacter changes)				_
MJ1532	1628155	1627745		34.1	68.9	219	
		64//201	prosphortbosyltormimno-5-aminoimidazole carboxamide ribotide isomerase {Methanococcus thermolithotrophicus}	51.9	8	411	
Biosynthesis	of cofactors,	prosthetic gr	Biosynthesis of cofactors, prosthetic groups, and carriers				
MJ0603	937289	938566	glutamate-1-cemialdehude aminotenacionalismos 1-semialdehude amino				
MJ0569	966316	067137	Supplier of the supplier of the subtility	51.7	70.6	1278	
207017	21.00	101107	porpnoollinogen deaminase (Bacillus subtilis)	41.2	61.4	822	
MJ0493	1035991	1036839	quinolinate phosphoribosyltransferase (Escherichia coli)	, 95			
MJ0407	1105699	1104965	quinolinate synthetase (Cyanonhora maradova)	5,5	0.10	849	
8411200	137761		(C) mobile paracova	37.2	58.8	735	
1000 1 100	136484	135309	S-adenosylhomocysteine hydrolase (Sulfolobus solfataricus)	617	70.5	1.37	
Biotin				;;;	(%)	0/11	
MJ1297	227704	177011					-7
	100	170/77	o-carboxynexanoate-CoA ligase (Bacillus sphaericus)	42.2	600	703	1-
MJ1298	227005	225890	8-amino-7-oxononanoate synthase (Bacillus enhagricus)		7.70	100	•
MJ1300	225025	223700		44.4	64.8	1116	
		223103	adenosylmethionine-8-amino-7-oxononanoate aminotransferase (Bacillus sphaericus)	39.9	64.2	1317	
MJ1619	1543130	1543552	bifunctional protein (Haemophilus influenzae)				
MJ1296	228286	228843		25.7	54.9	423	
		77077	olouin syntnetase (Bacillus sphaericus)	38.2	62.5	558	
M31299	225741	225100	dethiobiotin synthetase (Bacillus sphaericus)	37.0	29.0	643	
				1	7	7	

Heme and nornhamia	- Phinip						
200							
MJ1438	66330	65833	cobalamin (5'-phosphate) synthase (Escherichia coli)	26.1	48.7	408	
MJ0552	983686	984417	cobalamin biosynthesis J protein {Salmonella tyohimurium}	1 2			
MJ1314	212528	211842	Cobalamin biosynthesis protein D (Pseudomonas denitrificans)	70 6	717	132	
MJ0022	1448163	1447273	cobalamin biosynthesis protein D (Salmonella tynhimm:	78.0	o. ;	687	
MJ1569	1592308	1591700	cobalamin biosynthesis protein M (Salmonella tyrhimurium)	22.2	- :	168	
MJ1091	442661	443239	cobalamin biosynthesis protein M (Salmonella trattiamina)	2 3	24.7	609	
MJ0908	635150	631647	cobalamin biosynthesis protein N (Pseudomonas denitrificans)	23.7	74.4	579	
MJ0484	1046784	1045324	cobyric acid synthase {Methanococcus voltae}	2/5	o./5	3504	
MJ1421	85381	86352	Cobvrinic acid a c-diamide contrace (Columnation L.	/2:/	87.8	1461	
Minida	1333000	2,2001.1	Samonena typinimurium)	32.1	55.0	972	
CLICAL	1332000	1330965	glutamyl-tRNA reductase {Methanobacterium thermoautotrophicum}	47.8	6.99	1116	_
MJ0643	899800	898910	porphobilinogen synthase (Methanothermus sociabilis)	\$ 63	70.0	Γ	<u>72</u>
MJ0930	612059	611430	precorrin isomerase (Salmonella typhimurium)	18.7	6.67		_
MJ0771	780420	779932	precorrin-2 methyltransferase (Salmonella typhimurium)	30.4	55.0	480	
MJ0813	734876	735547	precorrin-3 methylase {Salmonella typhimurium}	44,	6.60	467	
MJ1578	1583277	1582501	precorrin-3 methylase {Salmonella typhimurium}	3 73	76.6	270	
MJ1522	1637017	1636385	preсотin-6Y methylase {Salmonella typhimurium}	20.6	5.5	///	PCT/
MJ0391	1116729	1117202	precorrin-8W decarboxylase (Salmonella tynhimurium)	2 2	3	500	US9
		1		73.9	49	474	7/

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MJ0965	573234	572509	uroporphyrin-III C-methyltransferase (Bacillus megaterium)	643	3 5		
MJ0994	549022	549444	uronombreinocen III and the time to the time	24.7	(7.2)	726	_
			archothings in synnase (bacillus subtilis)	27.8	49.4	423	
Menaquinone and ubiquinone	and ubiqui	попе					$\overline{}$
MJ1645	1509624	1508923	coenzyme PQQ synthesis protein III (Haemophilus influenzae)	13.3		907	
M lybdopterin				7776	23.3	707	_
MJ0824	725986	726762	molybdenum cofactor biosynthesis moad protein (Haemorahilus in fluoresis)	96,			
MJ0167	1301836	1302162	molybdenum cofactor biosymthesis mos B motoin (50.1.1.1.1.1.1.1.1.1.1.1.1.1.1.1.1.1.1.1	30.0	5/3		$\overline{}$
MII135	306360	. 105705	(Escribing coll)	46.4	9.69	327	_
	,,,,,,,	320/01	monybacenum cotactor biosynthesis moaC protein (Haemophilus influenzae)	49.2	70.9	423	_
MJ0886	654158	656017	molybdenum cofactor biosynthesis moeA protein {Escherichia coli}	372	5	0701	_
MJ0666	879771	880943	molybdenum cofactor biosynthesis moed profein / Haemonhilus in fl.	25.5	23.2	1800	
MJ1663	1491265	1490831	molyhdonterin enemine dinatoria Li	33.0	26.4	1173	 ;
100000	1000		Secretaring Secretaring United States of Synthesis protein A (Escherichia coli)	27.7	48.0	435	<u>.</u>
676 I CIAI	19////	197076	molybdopterin-guanine dinucleotide biosynthesis protein B (Escherichia coli)	32.2	577	55	_
Pantothenate						701	
MJ0913	626982	627779	pantothenate metabolism flavoprotein (Haemonhilus in flusaria)				
			- Charles IIII (Tracinopinius IIII (CHARLES)	77	, v	200	_

Riboflavin							_
MJ0055	1416688	1417278	GTP cyclohydrolase II {Bacillus subtilis}	36.9	66.0	103	
MJ0671	874773	875396	riboflavin-specific deaminase {Actinobacillus pleuronneumoniae}	5 5	2:0	160	
Thioredoxin, glutaredoxin, and glutathione	glutaredoxir	1, and glutath	1	25.5	65.5	974	
MJ1536	1622694	1623533	thioredoxin reductase {Mycoplasma genitalium}	38.6	8	970	
MJ0530	1005917	1005420	thioredoxin-2 {Saccharomyces cerevisiae}	3 5	0.00	940	
MJ0307	1184114	1184332	thioredoxin/glutaredoxin {Methanobacterium thermoautotronhicum }	23.0	6 8	498	
Thiamine)	09:5	419	
MJ1026	514172	515440	thiamine biosynthesis protein (Bacillus subrilis)		;		
M10601	040113	030400		45.0	99.1	1269	
	240113	939400	thiamine biosynthetic enzyme (Zea mays)	35.1	53.0	714	
Pyridine nucleotides	otides						74-
MJ1352	170567	171163	NH(3)-dependent NAD+ synthetase {Mycoplasma genitalium}	47.5	819	407	
			Cell envelope			166	
Membranes, lipoproteins, and porins	poproteins,	and porins					
MJ0544	508686	990443	dolichyl-phosphate mannose synthase {Trypanosoma brucei}	18.1	3 :	953	
MJ1057	475508	474981	glycosyl transferase (Neisseria gonorrhoeae)	35.5	1./5	650	
MJ0611	931098	930679	membrane protein (Socchamm en)	6.5.8	20.0	228	PCI
			Concording the latest	50.0	57.2	420	, 00
MJ0827	724322	723900	membrane protein (Homo sapiens)	077	0 63	423	

Murein sacculus and peptidoglycan	lus and pep	tidoglycan					٢
MJ1160	371691	370390	amidase (Moraxella catarrhalis)				_
MJ0204	1276277	1275219	amidonhoenhoilbonding	24.6	36.1	1302	_
Surface polysaccharides, linonolysacrhades	accharides.	linonolyeacel	erfee and anticon	52.0	72.9	1059	
7,00174							
4760CM	617598	618035	capsular polysaccharide biosynthesis protein (Staphylococcus aureus)	21.2	46.0	13.0	$\overline{}$
MJ1061	469649	470293	capsular polysaccharide biosynthesis profein D (Stark. 1		<u>Ş</u>	438	\neg
MJ1055	478643	477735	Compily lococcus aureus	56.3	72.2	645	
MILOSO		661111	capsular polysaccharide biosynthesis protein I (Staphylococcus aureus)	50.7	74.4	606	_
COLON	4/2220	471904	Capsular polysaccharide biosynthsis protein M (Staphylococcus aurene)	7.72	3	1	Т
MJ1607	1555624	1554455	LPS biosynthesis related rfbu-protein (Haemorphilus influenzae)		33.0	423	_
MJ1113	417528	418352	N. acad allocation in the state of the state	33.4	57.6	1170	_
710100				29.9	57.9	825	
MJUSSY	1110873	1112204	phosphomannomutase (Vibrio cholerae)	32.0	3		_
MJ1068	462901	464265	putative O-antigen transporter (Shinella games)	3/.0	27.8	1332	·クミ
MII066	464360	200777		24.5	46.6	1365	<u>-</u>
	404309	402430	spore coat polysaccharide biosynthesis protein C {Bacillus subtillis}	55.3	75.8	1963	_
MJ 1065	465444	466454	spore coat polysaccharide biosynthesis protein F (Bacillus subtillie)				<u> </u>
MJ1063	467331	467828	Shore cost notvesorheride himself.	37.9	59.0	101	
MIIOKO	050577		For section to broading specific (Bacillus subtillis)	36.0	55.4	498	
7001514	40/8/0	469279	spore coat polysaccharide biosynthesis protein G (Bacillus subtillis)	32.0	545	1410	_
MJ0211	1269601	1268732	UDP-glucose 4-epimerase (Streptococcus thermonhilus)				_
MJ1054	481027	478712	UDP-glucose dehydrogenase / Yanthomosas	35.1	24.8 8.	870	
M10428	1087466	100000	composition (variational campestris)	42.8	63.4	2316	
27,000	100/420	1088033	UDP-N-acetyl-D-mannosaminuronic acid dehydrogenase (Escherichia coli)	45.1	68.2	1200	
							_

Surface structures	tures						
MJ0891	650616	650005	flagellin B1 {Methanococcus voltae}	7 7 7	21.6	(1)	_
MJ0892	649880	649269	flagellin B2 {Methanococcus voltae}			710	_
MJ0893	649163	648516	flagellin B3 (Methanococcus voltag)	1.10	4.6.4	710	7
				59.1	78.7	648	
			Cellular processes				_
Cell division							7
MJ1489	10595	8721	cell division control protein {Saccharomyces cerevisiae}	24.0	63.3	250	_
MJ0363	1142460	1140220	cell division control protein 21 {Schizosaccharomyces nombe}	20.0	,,,,	18/5	T
MJ1156	375317	377947	cell division control protein CDC48 (Saccharomyces cerevision)	30.0	4.10	7741	_
MJ0169	1300988	1300329	cell division inhihitos (Dodine	51.9		2631	_
			Con civision minoriol (Dacillus subtillis)	28.8	51.2	099	_
MJ0579	957291	958088	cell division inhibitor (Bacillus subtillis)	31.8	512	708	76
MJ0547	988025	988732	cell division inhibitor {Bacillus subtillis}	32.8	22.7	96	-
MJ0084	1393471	1392869	cell division inhibitor minD (Escherichia coli)	32.1	50.4	807	_,_
MJ0174	1295971	1294976	cell division protein {Drosophila melanogaster}	78.7	20.4	500	-, -
MJ0370	1135876	1134956	cell division protein ftsZ {Anabaena 7120}	50.7	0.4.0	066	
MJ1376	147975	147343	cell division protein J {Haemophilus influenzae}	30.6	/11/	17,	
MJ0622	920029	921168	cell division protein Z {Hatoferax volcanii}	\$10	717	1140	_,_
MJ0148	1326798	1327538	centromere/microtubule-binding protein (Saccharomyces gerevisiae)	T		0711	
					04./	/4!	_

MJ1647	1508164	1507907	DNA binding protein (Methanococcus voltag)			
MJ1643	1513857	1510351	P115 protein (Myconlasma hyorhinic)	54.7	80.3	258
Chaperones				30.3	55.4	3507
MJ0999	543921	545471				
MJ0285	1202058	1202450	Chapteronin (internanopyrus kandleri)	73.5	87.6	1551
M10278	120226	6047071	near snock protein {Clostridium acetobutylicum}	29.0	44.6	402
0.1701	0/7/071	1207548	rotamase, peptidyl-prolyl cis-trans isomerase (Haemophilus influenzae)	, 6,	9 69	
MJ0825	725091	725765	rotamase, peptidyl-prolyl cie-frane feathers of December 1);	00.5	273
Detoxification			The second of th	31.8	8.09	675
MJ0736	804803	805753				
	00100	002423	alkyl hydroperoxide reductase (Sulfolobus solfataricus)	1.99	848	19
MJ1541	1618786	1619868	N-ethylammeline chlorohydrolase (Rhodococciis richananii		2	3
Protein and peptide secretion	ptide secre	tion	{snooped a notice of the control of	29.2	56.3	1083
MJ0478	1051985	1050570				77-
		9/90001	preprotein translocase secY (Methanococcus vannielii)	200	8 8 8	985
MJ0111	1365253	1364216	protein-export membrane protein (Strentomyces coelicales)	\dagger	90:0	1300
MJ1253	276673	277377	Protein-export membrane arctain (E. L.: 1:	25.9	51.7	1038
MJ0260	1226000	1133661		30.5	57.0	705
	1220030	1220044	signal peptidase {Canis familiaris}	326	2 7 5	l ä
MJ0101	1376106	1377308	signal recognition particle protein (Haemonhilus influenzae)	十		3
MJ0291	1198470	1197244	Signal recognition marticle martinities of the signal recognition and the signal signa	42.0	9.19	1203
				48.3	69.4	1227

Transformation	uc						_
MJ0781	768702	770798	klbA protein (Plasmid RK2)	34.6	540	2007	_
MJ0940	602402	601929	transformation sensitive protein {Homo sapiens}	35.0	53.0	474	
Cellular processes	sses						_
MJECL17	20110	19889	archaeal histone (Pyrococcus sp.)	58.8	0.18	221	
MJECL29	36456	26220	archaeal histone (Pyrococcus sp.)	64.2	83.6	236	
MJ1258	271686	271486	archaeal histone (Pyrococcus sp.)	71.7	83.6	201	
MJ0168	1301348	1301548	archaeal histone (Pyrococcus sp.)	67.2	86.6	201	
MJ0932	610153	609953	archaeal histone {Pyrococcus sp.}	67.2	86.6	201	
			Central intermediary metabolism				-7
Amino sugars							78-
MJ1420	90244	86939	glutaminefructose-6-phosphate transaminase {Escherichia coli}	41.2	818	3306	
Degradation of polysaccharides	polysaccha	ırides				2000	
MJ1611	9180551	1549542	alpha-amylase (Pyrococcus furiosus)	27.0	\$ 0.5	1775	
MJ0555	981500	980529	endoglucanase (Homo sapiens)	1 44	8 99	977	
MJ1610	1551992	1550967	glucoamylase (Clostridium sp)	28.0	49.2	1026	

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Other							1
MJ1656	1498675	1497965	2-hydroxyhepta-2 4-diene-1 7-diosta incensor (r	-			3007
MJ0406	1106800	1105907	ribokinaca (Echaciatic att)	40.2	9.19	711	a3U
MJ0309	1182260	1.0.011		23.2	46.3	894	
Dhara	(778)1	11830//	urechydrolase (Methanothermus fervidus)	40.9	60.7	819	
r nospaoras compounds	spunoduo						_
MJ0963	575418	577049	N-methylhydantoinase (Arthrohacter en)				
MJ0964	573516	575145	Namehilling.	32.6	53.0	1632	
Polyamine biosynthesis	osynthesis		14-mem mydantomase {Arthrobacter sp.}	37.7	56.4	1830	· · · · ·
MJ0535	1001006	100001					
	2001001	1002031	acetylpolyamine aminohydolase (D01044 Mycoplana)	11.1	787	1007	
MJ0313	1179250	1179801	Spermidine synthase (Homo sapiens)		200	1020	
Polysaccharides-(cytoplasmic)	es-(cytoplas	mic)		32.3	57.7	552	-7
MIIGOR	1666969						9-
2001211	1222626	155/354	glycogen synthase (Hordeum vulgare)	3 5			
Nitrogen metabolism	bolism			133.7	58.3	1497	
MJ1187	345237	344335	ADP-ribosylg/ycohydrolaes (des) (Dt. 1.1.				_•
MJ0713	824113	826278	hydrogenace accessors accessors to the control of t	29.8	50.8	903	
MJ0214	1267658	1267214	AZOIOOACIEL CHIOOCOCCUM)	33.8	54.8	2166	
MIDEZE		F15/071	nydrogenase accessory protein {Azotobacter chroococcum}	30.7	56.5	345	P
0/000701	869511	870276	hydrogenase expression/formation protein {Rhizobium leguminesarum}				CT
MJ0442	1075480	1076028	hydrogenase expression/formation profein R / Rhizakim.	40.	65.3	996	/US9
MJ0200	1279494	1279739	hydropenase expression/6	44.6	64.0	549	7/14
			"Jaioballase capiession formation protein C {Azotobacter vinelandii}	40.0	68.8	246	900

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MJ0993	549539	550525	hydrogenase expression/formation protein D {Alcaligenes eutrophus}	44.7	63.5	987
MJ0631	914544	914089	hydrogenase maturation protease {Escherichia coli}	33.9	58.9	456
MJ1093	441468	440584	nifB protein {Anabaena sp}	43.1	67.2	885
MJ0879	667622	666984	nitrogenase reductase (Methanococcus voltae)	77.2	89.1	639
MJ0685	859442	858696	nitrogenase reductase related protein (Clostridium pasteurianum)	31.7	49.6	747
MJ1051	483344	484411	nodulation factor production protein (Bradyrhizobium japonicum)	32.1	51.1	1068
MJ1058	473947	473141	nodulation factor production protein (Bradyrhizobium japonicum)	37.7	58.0	807
Carbon Fixation	0.0					
MJ0152	1325036	1322820	carbon monoxide dehydrogenase, alpha subunit (Clostridium thermoaceticum)	42.1	9.59	2217
MJ0153	1322553	1320256	carbon monoxide dehydrogenase, alpha subunit (Methanothrix soehngenii)	47.0	67.3	270g
MJ0156	1319256	1317883	carbon monoxide dehydrogenase, alpha subunit {Clostridium thermoaceticum}	47.8	\$ 69	1374
MJ0728	809951	811783	carbon monoxide dehydrogenase, beta subunit (Rhodospirillum rubrum)	35.9	55.0	
MJ0112	1362285	1363667	corrinoid/iron-sulfur protein, large subunit {Clostridium thermoaceticum}	32.9	55.1	1383
MJ0113	1361128	1362030	corrinoid/iron-sulfur protein, small subunit {Clostridium thermoaceticum}	37.7	58.8	903
MJ1235	292453	293673	ribulose bisphosphate carboxylase, large subunit (Synechococcus sp)	\top	60.3	1221
				_	-	

			Energy metabolism			
Aerobic						
MJ0649	896262	894919	NADH oxidase {Enterococcus faecalis}			
MJ0520	1011104	1011892	() () () () () () () () () ()	0.62	50.4	1344
Anserobic				29.5	53.9	789
M10092	1395740	1304303				
700000	1303/40	1364282	umarate reductase {Thermoplasma acidophilum}	40.2	57.0	1467
ATP-proton motive force interconversion	notive force	interconversi	U(
MJ0217	1263468	1265171	ATP synthase, subunit A {Enterococcus hirae}	6.9	1	
MJ0216	1265356	1266615	1100	5.00	0.0	1/04
MJ0219	1261985	1263040		9.4	84.5	1260
MIDELS	006134	277700	Security (Transletax Volcanii)	28.1	50.0	1056
CIOOCIAI	920124	926663	ATP synthase, subunit D (Enterococcus hirae)	34.8	8.99	540
MJ0220	1261297	1261737	ATP synthase, subunit E {Methanosarcina mazeii}	600	000	
MJ0218	1263054	1263347		1		1 2
MJ0222	1258252	1260294		7	1.72	294
MJ0221	1260641	1261060		十	2775	2043
				34.6	59.8	420

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Electron transport	Insport						r
MJ1446	57416	56646	cytochrome-c3 hydrogeness games 11:100				
MJ0741	803000	803320	desulfoferrodovia (Dames de la	40.1	52.4	171	
MJ0578	058004	06000	Complete Control (Destinovibrio Vulgaris)	44.0	59.4	321	
MISSE	10000	┿	lemedoxin (Clostridium sticklandii)	49.1	56.9	807	_
1 ODDCIMI	1411998	1411759	ferredoxin (Methanococcus thermolithotrophicus)				
MJ0722	815808	816038	ferredoxin (Methanobacterium thermonistration)	42.9	29.0	240	
MJ0099	1379076	1379456	ferredoxin (Desultovibrio dominio de la companya de	42.3	9.09	231	
MJ0199	12700761	+	described describing describing as a second	40.0	62.0	381	
M10633	0/66/21		terredoxin {Methanococcus thermolithotrophicus}	74.6	84.8	186	
1410233	1003408	1003575	ferredoxin 2[4Fe-4S] homolog (Methanosarcina thermorbile)				
MJ0624	918981	918808	ferredoxin 2[4Fe=4S] {Methanosarcina shormontile;	36.9	54.4	89	
MJ0267	1217567	1218463	Series de la company de la com	48.0	68.0	174	-8
		1210403	refredoxin oxidoreductase, alpha subunit (Klebsiella pneumoniae)	20.4	ç		2-
MJ0276	1209645	1210727	ferredoxin oxidoreductase alpha enhimit (Unitable)	*:/3	20.7	89/	•
MJ0266	1218644	1219387	ferration:	44.5	63.0	1083	
Miness	2000		Concedent oxidoreductase, beta subunit (Klebsiella pneumoniae)	32.6	51.0	744	
/ Contai	398693	999424	ferredoxin oxidoreductase, beta subunit (Halobacterium halobium)];			
MJ0268	1217015	1217272	ferredoxin oxidoreductase, delta subunit (Dunggonn 6;	?	-:-	132	
MJ0536	999441	086666	ferredoxin oxidoreductors	58.9	71.8	258	
MJ0269	1216601	1216903	Service Strong Countries, gamma subunit (Pyrococcus furiosus)	32.0	50.9	540	
M10732	020200	001000	refredoxin oxidoreductase, gamma subunit (Pyrococcus furiosus)	55.6	74.7	393	
	0/6000	808100	flavoprotein (Methanobacterium thermoautotrophicum)	40 4	2 5	121	
MJ1192	339066	338095	methylviologen-reducing hydrogenase, alpha chain {Methanococcus voltae}	†	+		
					66.0	9/2	

MJ1191	340221	339385	methylviologen-reducing hydrogenase, gamma chain (Methanococcus voltae)	71.5	63.3	03.7
MJ1362	160414	161055	NADH dehydrogenase, subunit I (Mitochondrion Oncorhynchus)	; ;	5.50) 50
MJ0514	1016474	1017223	polyferredoxin (Methanococciie waltan)	1.63	0.00	047
M 10034			Control of the contro	36.7	52.5	750
MJ0934	608147	607521	polyferredoxin (Methanothermus fervidus)	40.9	54.3	627
MJ1303	220214	221701	polyferredoxin (Methanobacterium thermoautotrophicum)	30 \$	1 75	1,400
MJ1193	337655	336591	polyferredoxin (Methanococcus voltae)	27.7	100.	0041
MJ1227	301853	301257	pyruvate formate-lyase activating enzyme (Clostridium nacteurismum)	, , ,		2001
MJ0735	805546	805785	rubredoxin (Clostridium thermospherelistics)	51.4	20.0	297
				59.7	77.0	240
MJ0740	803522	803659	rubredoxin {Clostridium thermosaccharolyticum}	3 79	2 70	130
Fermentation				22.5	04.0	001
MJ0007	1463447	1462359	2-hydroxyglutaryl-CoA dehydratase, subunit beta (Acidaminococcus	22.6	48.2	6801
Gluc neogenesis	si					
MJ1479	22527	21358	alanine aminotransferase 2 {Panicum miliaceum}	30.1	0 0	97.
MJ0542	991264	994794	phosphoenolpyruvate synthase (Pyrococcus furiosus)	100	2 6	2/1
				700	×	1531

Glyc lysis							ſ
MJ1482	18946	18044	2				
MIDEAT	200.00		4-prosphoglycerate kinase (Methanothermus fervidus)	47.1	70.9	903	<u> </u>
1400011	901393	902325	3-phosphoglycerate kinase (Methanothermus fervidus)				Т
MJ0232	1248239	1249432	enolase (Bacillus subtilis)	28.2	78.1	933	- T
MJ1605	1557395	1558597	Oliver 6 about 1	57.7	78.2	1194	_
M11146	102001		siccose-o-pinospilate isomerase (Bacillus stearothermophilus)	32.3	54.6	1203	
2	280093	38/055	glyceraldehyde 3-phosphate dehydrogenase (Methanothermus fervidus)	205	3,50	3	Т
MJ0490	1038560	1037697	lactate dehydrogenase (Thermotoga maritima)	2,2	?	E P	-
MJ1411	100555	65100	January 19	39.9	63.2	864	
	CCC001	79167	NADP-dependent glyceraldehyde-3-phosphate dehydrogenase (L15191 Streptococcus)	39.2	59.6	1389	т—
MJ0108	1367951	1366716	Dynivate kinace (Bacillia state)				,
MIICO	100.00		commend because scaroinesmobilities	39.1	60.5	1236	
07C1Clai	1631071	1631589	triosephosphate isomerase (Mycoplasma genitalium)	38.5			_
Pentose phosphate pathway	bate pathwa	'n		79.0	49.1	\$19	-8
MIDERO	707370	9//003					4.
	902404	800083	pentose-5-phosphate-3-epimerase (Solanum tuberosum)	44.5	3 5	3	-
MJ1603	1560724	1560047	ribose 5-phosphate isomerase (Mus musculus)	,	07.7	000	
MJ0960	580121	A52082		42.0	63.4	878	
		2/22/	uansaruojase { Bacillus subtilis}	60.7	79.5	456	_
MJ0681	864603	865355	transketolase' (Homo sapiens)			3	
MJ0679	866375	867073	fransketolese" (Homo conicas)	43.7	28.5	753	
			(Tollio sapiens)	36.0	61.3	669	
							_

Pyruvate dehydrogenase	hydrogenase					
MJ0636	906464	006202	71.			
Sugars		767606	unydroupoamide dehydrogenase (Haloferax volcanii)	28.9	51.0	1173
MJ1418	91211	69906	ficulate 1 whom to 11			
TCA cycle				29.1	48.7	543
MJ0499	1031331	1032530	aconitace / Conchast			
MJ1294	229770	230381	firmancia (Saccinarom) yes cerevisiae)	29.7	49.8	1200
MJ0617	925239	974778	G	35.1	55.7	612
MJ1596	1568967	1569908	innarate nydratase, class I " (Bacillus stearothermophilus)	43.8	9.99	462
MJ0720	817433	818431	isociuate denydrogenase {Thermus aquaticus}	42.9	61.4	1032
MJ1425	77051	76299	malate dehydrogenase (NADP) {Thermus aquaticus}	48.0	64.7	666
MJ0033	1438609	1437116		61.3	77.6	753
MJ1246	282664	283449	SuccinvI-CoA everthetees ItaVoprotein subunit {Escherichia coli}	41.8	58.1	1494
MJ0210	1271318	1270227	succinal Coverage, alpha subunit (Escherichia coli)	9.65	74.8	786
			succinyi-CoA syntnetase, beta subunit (Thermus aquaticus)	48.8	68.7	1092

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_	_		

Meth	1					
Membrogenesis	515					
MJ0253	1232773	1232405	8-hydroxy-5-deazaflavin-reducing hydrogenase, delta subunit {Methanobacterium thermoautotrophicum}	47.1	0.17	698
MJ1035	505234	506022	coenzyme F420-dependent NS,N10-methylene-tetrahydromethanopterin dehydrogenase {Methanobacterium thermoautotrophicum}	66.5	79.8	789
MJ0727	811895	812725	coenzyme F420-reducing hydrogenase, alpha subunit {Methanobacterium thermoautotrophicum}	26.8	45.8	831
MJ0029	1442517	1441279	coenzyme F420-reducing hydrogenase, alpha subunit (Methanococcus voltae)	50.3	1.99	1239
MJ0030	1441022	1440558	coenzyme F420-reducing hydrogenase, alpha subunit (Methanococcus voltae)	66.5	83.3	465
MJ1349	175566	176222	coenzyme F420-reducing hydrogenase, beta subunit {Methanococcus voltae}	36.6	55.7	657
MJ0725	813779	814453	coenzyme F420-reducing hydrogenase, beta subunit {Methanobacterium thermoautotrophicum}	41.0	62.0	675
MJ0870	677657	679372	coenzyme F420-reducing hydrogenase, beta subunit (Methanobacterium thermoautotrophicum)	42.7	63.2	1716
MJ0032	1439835	1438990	coenzyme F420-reducing hydrogenase, beta subunit (Methanococcus voltae)	72.0	85.5	846
MJ0726	812987	813499	coenzyme F420-reducing hydrogenase, gamma subunit {Methanococcus voltae}	42.7	59.4	513
MJ0031	1440505	1439873	coenzyme F420-reducing hydrogenase, gamma subunit {Methanococcus voltae}	75.5	87.3	633
MJ0295	1192687	1193304	formate dehydrogenase (fdhD) (Wolinella succinogenes)	35.6	57.7	618
MJ0006	1463887	1465020	formate dehydrogenase, alpha subunit (Methanobacterium formicicum)	41.6	61.1	1134
MJ1353	168767	170344	formate dehydrogenase, alpha subunit (Methanobacterium formicicum)	54.2	6.07	1578
MJ0005	1465405	1466247	formate dehydrogenase, beta subunit {Methanobacterium formicicum}	49.5	72.1	843

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MJ0155	1319767	1319315	formate dehydrogenase, iron-sulfur subunit (Wolinella surcingenes)	.;	3		_
MJ0264	1220122	1220433	formate hydrogenlyase, subunit 2 (Escherichia coli)	41.7	86.9	453	
MJ0265	1219502	1219930	formate hydrogenivase submit 2 (Becherickie ett.)	42.9	865	312	
MJ0515	1013710	1014735	formate hydrogen by the contract of the contra	45.5	0.19	429	
MJ1027	514001	512871	formate hydrogen trace anti-it for the	31.0	51.1	1026	
MJ1363	159614	160018	Security and secur	34.3	53.3	1131	
MIOSIA	1012167	91000	Jorniare nydrogenlyase, subunit 7 {Escherichia coli}	38.4	60.9	405	
110318	/cicioi	1013600	formate hydrogenlyase, subunit 7 (Escherichia coli)	48.8	65.6	444	
MJU518	1175065	1175823	formylmethanofuran:tetrahydromethanopterin formyltransferase (Methanobacterium thermoautotrophicum)	68.6	84.5	759	
MJ1338	185930	185007	H(2)-dependent methylenetetrahydromethanopterin dehydrogenase related protein {Methanobacterium thermoautotrophicum}	29.1	50.5	924	
MJ0715	823334	822423	H2-forming N5,N10-methylene-tetrahydromethanopterin dehydrogenase-related protein {Methanococcus voltae}	29.9	52.5	912	-
MJ0784	765279	764272	tetrahydromethanopterin dehydrogenease	73.6	85.5	8001	87-
MJ1190	342199	341003	Suhinit A Mothant	7			
MJ0743	801736	802422	+	28.0	75.2	1197	
MJ0863	684944	685798	+	59.3	79.0	687	•
MJ0744	801103	801489	Sulfide reductase cubunit (Machanical Control	63.2	80.2	855	
MJ0864	684283	684840	sulfide reduction of the control of	53.4	68.4	387	1 66 0
MJ0118	1357167	135667	(internal thermosulotrophicum)	52.6	669	558	147
			Contains in reductase II operon, protein D (Methanothermus fervidus)	53.2	77.5	501	J U

MJ0083	1395319	1393880	methyl commune Mandace II all the				Г
10001	2022061			868	95.5	1440	
18000[M	1397/00	1396351	methyl coenzyme M reductase II, beta subunit {Methanothermus fervidus}	79.7	89.4	1350	
MJ0082	1396335	1395538	methyl coenzyme M reductase II, gamma subunit {Methanothernus fervidus}	83.0	92.1	798	
MJ0844	702037	701465	methyl coenzyme M reductase operon, protein C {Methanococcus vannielii}	82.5	926	\$72	_
MJ0843	702395	702069	methyl coenzyme M reductase operon, protein D (Methanococcus voltae)	58.0	814	127	
MJ1662	1491537	1493201	methyl coenzyme M reductase system, component A2 {Methanobacterium thermoautotrophicum}	37.1	1.09	1665	
MJ1242	284878	286338	methyl coenzyme M reductase system, component A2 {Methanobacterium thermoautotrophicum}	6.09	77.8	1461	
MJ0846	700322	698880	methyl coenzyme M reductase, alpha subunit (Methanococcus voltae)	86.1	92.1	1443	
MJ0842	703907	702576	methyl coenzyme M reductase, beta subunit {Methanococcus vannielii}	75.3	87.4	1332	
MJ0845	701389	700673	methyl coenzyme M reductase, gamma subunit {Methanococcus yannielii}	78.7	6 2	717	
MJ1636	1520054	1519128	N5,N10-methenyl-tetrahydromethanopterin cyclohydrolase {Methanobacterium thermoautotrophicum}	9.69	82.3	927	-88-
MJ1534	1625526	1624534	N5,N10-methylene tetrahydromethanopterin reductase {Methanobacterium thermoautotrophicum}	66.2	79.7	993	
MJ0850	696203	695895	N5-methyl-tetrahydromethanopterin:coenzyme M methyltransferase {Methanobacterium thermoautotrophicum}	36.6	59.8	309	
MJ0849	696884	696216	N5-methyl-tetrahydromethanopterin:coenzyme M methyltransferase (Methanobacterium thermoautotrophicum)	41.8	62.3	699	CT/US:
MJ0852	695117	694914	N5-methyl-tetrahydromethanopterin:coenzyme M methyltransferase {Methanobacterium thermoautotrophicum}	37.1	64.6	204	7//14900
				-			<u>'</u>

MJ0851 695866 695138 NS-n MJ0847 698519 697749 NS-n MJ0854 694607 693651 NS-n MJ0848 697696 697043 NS-n MJ0853 694857 694639 NS-n MJ1169 363822 362122 tungs MJ1194 336096 335260 tungs MJ1171 361740 360973 tungs	16.mothy totrohydromethene				
698519 697749 694607 693651 697696 697043 694857 694639 336322 362122 336366 335260	(Methanobacterium thermoautotrophicum)	55.2	73.5	729	98/0/
694607 693651 697696 697043 694857 694639 363822 362122 336096 335260 361740 360973	N5-methyl-tetrahydromethanopterin:coenzyme M methyltransferase (Methanobacterium thermoautotrophicum)	58.3	76.4	177	
697696 697043 694857 694639 363822 362122 336096 335260 361740 360973	N5-methyl-tetrahydromethanopterin:coenzyme M methyltransferase {Methanobacterium thermoautotrophicum}	62.1	77.5	957	
363827 694639 363822 362122 336096 335260 361740 360973	NS-methyl-tetrahydromethanopterin:coenzyme M methyltransferase {Methanobacterium thermoautotrophicum}	63.5	77.8	654	
363822 362122 336096 335260 361740 360973	N5-methyl-tetrahydromethanopterin:coenzyme M methyltransferase G {Methanobacterium thermoautotrophicum}	51.1	76.6	219	
336096 335260 361740 360973	tungsten formylmethanofuran dehydrogenase, subunit A {Methanobacterium thermoautotrophicum}	69.4	81.5	1701	т—
361740 360973	tungsten formylmethanofuran dehydrogenase, subunit B {Methanobacterium thermoautotrophicum}	71.1	84.0	837	
therm	tungsten formylmethanofuran dehydrogenase, subunit C {Methanobacterium thermoautotrophicum}	52.7	67.7	768	89-
MJ0658 887575 886886 tungst	tungsten formylmethanofuran dehydrogenase, subunit C related protein {Methanobacterium thermoautotrophicum}	35.4	53.4	069	
MJ1168 364202 363852 tungst	tungsten formylmethanofuran dehydrogenase, subunit D {Methanobacterium thermoautotrophicum}	55.2	74.8	351	·
MJ1165 366038 365637 tungst	tungsten formylmethanofuran dehydrogenase, subunit E {Methanobacterium thermoautotrophicum}	38.3	61.1	402	PCT/US
MJ1166 365484 364567 tungster	tungsten formylmethanofuran dehydrogenase, subunit F (Methanobacterium thermoautotrophicum)	47.6	67.4	918	-T

MJ1167	364516	364271	tungsten formylmethanofuran dehydrogenase, subunit G {Methanobacterium thermoautotrophicum}	43.1	58.5	246	
Fatty acid and phospholipid metabolism	d phospholi	pid metabolis		_			T
MJ0705	840072	838927	3-hydroxy-3-methylelutaryl coenzyme A reduction (11)				Т
MJ1546	1612371	1611697	acyl carrier protein synthese (Pyroposis 6	49.8	67.3	1146	Т
MJ0860	969889	689499	hifurctional chock their in the contract of the contract of the chock their in the contract of the chock their interest of the chock the	63.1	78.0	675	
			thermoautotrophicum}	49.5	71.7	804	
MJ1229	299478	300644	biotin carboxylase {Anabaena sn}				Т
MJ1212	316220	316792		58.9	76.2	1167	
	210257	210/00	CDF-diacylglycerol-serine O-phosphatidyltransferase {Bacillus subtilis}	45.5	63.7	558	1
MJ1504	1661217	1662188	lipopolysaccharide biosynthesis protein (bpID) (Bordetella pertussis)	44.2	3	6	T-
MJ1087	446091	445231	melvalonate kinase (Schizosaccharomyces nomba)		1.5	7//2	7
MJ1549	1610772	1609735	noncomposition limit demonstrate and positions from the positions and the positions are provided the positions and the positions are provided to the positions and the positions are provided to the position are provided to the positions are provided to the position are provided to the positions are provided to the positions are provided to the positions are provided to the position are provided to the position are provided to the position are provided t	31.5	53.7	861	\neg
			indiappearing inplu-transfer protein {Pyrococcus furiosus}	46.9	0.99	1038	90
			Purines, pyrimidines, nucleosides, and nucleotides				<u>}~</u>
2'-Deoxyribonucleotide metabolism	ucleotide m	etabolism					_
MJ0832	719820	714604	angerobic ribonucleoside triphombat				-
MJ0430	1085407	1002000	conscionation of the second coling to the second coling co	28.1	49.9	5217	
	1620001	6000001	deoxycytidine triphosphate deaminase (Desulfurolobus ambivalens)	40.4	5.19	513	_
MJI 102	429115	428648	deoxycytidine triphosphate deaminase, putative (Desulfurolobus amhivalene)	3.5	1		_
MJ0511	1019410	1020075	deoxyuridylate hydroxymethylase (Methanohacterium the	34.1	23.2	468	-
M10937	696963	504031	(moindonnaidh meimeannaidh mainneannaidh an	39.4	59.6	999	
	2000	004721	Blycinamide ribonucleotide synthetase {Homo sapiens}				_

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42.6 67.4 41.0 59.1 41.4 66.7 52.0 72.3 52.0 72.3 51.5 68.3 61.8 80.4 51.5 68.3 61.8 80.4 51.5 68.3 62.1 filis} 52.9 71.5 fis} 61.8 75.9 fis}								
613484 612135 adenylosuccinate lyase {Bacillus subtilis} 42.6 67.4 976592 975741 adenylosuccinate synthetase {Haemophilus influenzae} 41.0 59.1 5 1586386 1585823 GMP synthetase {Borrelia burgdorferi} 41.4 66.7 399509 400264 GMP synthetase {Haemophilus influenzae} 52.0 72.3 5 154271 inosine-S-monophosphate dehydrogenase {Pyrococcus furiosus} 61.8 80.4 6 262116 262436 nucleoside diphosphate kinase {Haemophilus influenzae} 56.3 76.2 7 262116 262436 nucleoside diphosphate kinase {Haemophilus influenzae} 31.5 68.3 8 925486 925941 phosphoribosylformylglycinamidine cyclo-ligase {Bacillus subtilis} 56.3 76.2 1277597 1276734 phosphoribosylformylglycinamidine synthase I {Bacillus subtilis} 52.9 71.5 262585 264714 phosphoribosylformylglycinamidie formyltransferase 2 {Bacillus subtilis} 43.3 65.1 13611 14633 phosphoribosylglycinamide formyltransferase 2 {Bacillus subtilis}	Purine ribon	ucleotide bio	synthesis .					_
976592 975741 adenylosuccinate synthetase (Haemophilus influenzae) 42.0 07.4 1586386 1585823 GMP synthetase (Borrelia burgdorferi) 41.4 66.7 399509 400264 GMP synthetase (Haemophilus influenzae) 52.0 72.3 1545605 1544271 inosine-5-monophosphate dehydrogenase (Pyrococcus furiosus) 61.8 80.4 262116 262436 nucleoside diphosphate kinase (Haemophilus influenzae) 51.5 68.3 925486 925941 phosphoribosylaminoimidazole carboxylase (Methanobrevibacter smithii) 56.3 76.2 12772482 1572009 phosphoribosylformylglycinamidine cyclo-ligase (Bacillus subtilis) 51.0 69.1 1277597 1276734 phosphoribosylformylglycinamidine synthase II (Bacillus subtilis) 52.9 71.5 262585 264714 phosphoribosylglycinamidie formyltransferase 2 (Bacillus subtilis) 61.8 75.9 13611 14633 phosphoribosylglycinamide formyltransferase 2 (Bacillus subtilis) 61.8 75.9 155580 156431 ribose-phosphate pyrophosphokinase (Haemophilus influenzae) 71.5 66.1 <td>MJ0929</td> <td>613484</td> <td>612135</td> <td>adenylosuccinate lyase {Bacillus subtilis}</td> <td>7,4</td> <td></td> <td></td> <td>-</td>	MJ0929	613484	612135	adenylosuccinate lyase {Bacillus subtilis}	7,4			-
1586386 1585823 GMP synthetase {Haemophilus influenzae} 41.0 59.1 1586386 1585823 GMP synthetase {Borrelia burgdorferi} 41.4 66.7 159509 400264 GMP synthetase {Haemophilus influenzae} 52.0 72.3 1545605 154271 inosine-5'-monophosphate dehydrogenase {Pyrococcus furiosus} 61.8 80.4 262116 262436 nucleoside diphosphate kinase {Haemophilus influenzae} 51.5 68.3 925486 925941 phosphoribosylaminoimidazole carboxylase {Methanobrevibacter smithii} 56.3 76.2 1572482 1572009 phosphoribosylformylglycinamidine cyclo-ligase {Bacillus subtilis} 51.0 69.1 1507541 1507071 phosphoribosylformylglycinamidine synthase II {Bacillus subtilis} 52.9 71.5 13611 14633 phosphoribosylglycinamide formyltransferase 2 {Bacillus subtilis} 61.8 75.9 155580 156431 ribose-phosphokinase {Haemophilus influenzae} 41.1 65.5 155580 156431 ribose-phosphokinase {Haemophilus influenzae} 41.1 65.5 155580 156431 ribose-phosphokinase {Haemophilus influenzae} 61.8 75.9 15658 156491 ribose-phosphokinase {Haemophilus influenzae} 75.9 15658 156991	MJ0561	976597	975741		47.0	67.4	1350	T
1586386 1585823 GMP synthetase {Borrelia burgdorferi}} 41.4 66.7 399509 400264 GMP synthetase {Haemophilus influenzae}} 52.0 72.3 1545605 1544271 inosine-5'-monophosphate dehydrogenase {Pyrococcus furiosus} 61.8 80.4 262116 262436 nucleoside diphosphate kinase {Haemophilus influenzae} 51.5 68.3 925486 925941 phosphoribosylaminoimidazole carboxylase {Methanobrevibacter smithii} 56.3 76.2 1572482 1572009 phosphoribosylformylglycinamidine cyclo-ligase {Bacillus subtilis} 51.0 69.1 1277597 1276734 phosphoribosylformylglycinamidine synthase I {Bacillus subtilis} 52.9 71.5 262585 264714 phosphoribosylformylglycinamide formyltransferase 2 {Bacillus subtilis} 61.8 75.9 13611 14633 phosphoribosylglycinamide formyltransferase 2 {Bacillus subtilis} 61.8 75.9 155580 156431 ribose-phosphate pyrophosphokinase {Haemophilus influenzae} 74.1 65.5			1010	adenyiosuccinate synthetase (Haemophilus influenzae)	41.0	59.1	852	
399509 400264 GMP synthetase {Haemophilus influenzae} 52.0 72.3 1545605 1544271 inosine-5'-monophosphate dehydrogenase {Pyrococcus furiosus} 61.8 80.4 262116 262436 nucleoside diphosphate kinase {Haemophilus influenzae} 51.5 68.3 925486 925941 phosphoribosylaminoimidazole carboxylase {Methanobrevibacter smithii} 56.3 76.2 1572482 157209 phosphoribosylaminoimidazolesuccinocarboxamide synthase {Bacillus subtilis} 51.0 69.1 1277597 1276734 phosphoribosylformylglycinamidine synthase I {Bacillus subtilis} 52.9 71.5 262585 264714 phosphoribosylformylglycinamide formyltransferase 2 {Bacillus subtilis} 61.8 75.9 13611 14633 phosphoribosylglycinamide formyltransferase 2 {Bacillus subtilis} 61.8 75.9	MJ1575	1586386	1585823	GMP synthetase {Borrelia burgdorferi}	414	647	793	_
1545605 1544271 inosine-5'-monophosphate dehydrogenase {Pyrococcus furiosus} 61.8 80.4 262116 262436 nucleoside diphosphate kinase {Haemophilus influenzae} 51.5 68.3 925486 925941 phosphoribosylaminoimidazole carboxylase {Methanobrevibacter smithii} 56.3 76.2 1572482 1572009 phosphoribosylaminoimidazole carboxylase {Methanobrevibacter smithii} 56.3 76.2 1277597 1276734 phosphoribosylformylglycinamidine cyclo-ligase {Bacillus subtilis} 42.7 64.4 1507541 1507071 phosphoribosylformylglycinamidine synthase II {Bacillus subtilis} 52.9 71.5 262585 264714 phosphoribosylformylglycinamidine synthase II {Bacillus subtilis} 61.8 75.9 13611 14633 phosphoribosylglycinamide formyltransferase 2 {Bacillus subtilis} 34.1 64.8 155580 156431 ribose-phosphate pyrophosphokinase {Haemophilus influenzae} 34.1 64.8 64.8 155580 156431 ribose-phosphate pyrophosphokinase {Haemophilus influenzae} 34.1 64.8 64.	MJ1131	399509	400264	GMP synthetase {Haemophilus influenzae}	5	35.	, oc	-,-
262116 262436 nucleoside diphosphate kinase {Haemophilus influenzae} 51.5 68.3 925486 925941 phosphoribosylaminoimidazole carboxylase {Methanobrevibacter smithii} 56.3 76.2 1572482 1572009 phosphoribosylaminoimidazolesuccinocarboxamide synthase {Bacillus subtilis} 51.0 69.1 1277597 1276734 phosphoribosylformylglycinamidine cyclo-ligase {Bacillus subtilis} 52.9 71.5 1507541 1507071 phosphoribosylformylglycinamidine synthase II {Bacillus subtilis} 52.9 71.5 262585 264714 phosphoribosylformylglycinamidine synthase II {Bacillus subtilis} 65.1 75.9 13611 14633 phosphoribosylgycinamide formyltransferase 2 {Bacillus subtilis} 61.8 75.9 155580 156431 ribose-phosphate pyrophosphokinase {Haemophilus influenzae} 34.1 65.5	MJ1616	1545605	1544271	inosine-5'-mononhosenhote detrudenment	22.0	77.3	126	_
262116 262436 nucleoside diphosphate kinase {Haemophilus influenzae} 51.5 68.3	7701174				8.19	80.4	1335	
925486925941phosphoribosylaminoimidazole carboxylase (Methanobrevibacter smithil)\$6.376.215724821572009phosphoribosylaminoimidazolesuccinocarboxamide synthase (Bacillus subtilis)\$1.069.112775971276734phosphoribosylformylglycinamidine cyclo-ligase (Bacillus subtilis)\$2.971.5262585264714phosphoribosylformylglycinamidine synthase II (Bacillus subtilis)\$43.365.11361114633phosphoribosylglycinamide formyltransferase 2 (Bacillus subtilis)\$41.365.1155580156431ribose-phosphate pyrophokinase (Haemophilus influenzae)\$41.5\$4.5	MJ1263	262116	262436	nucleoside diphosphate kinase (Haemophilus influenzae)	515	68.3	22.1	_
15724821572099phosphoribosylaminoimidazolesuccinocarboxamide synthase {Bacillus subtilis}51.069.112775971276734phosphoribosylformylglycinamidine cyclo-ligase {Bacillus subtilis}42.764.415075411507071phosphoribosylformylglycinamidine synthase II {Bacillus subtilis}52.971.5262585264714phosphoribosylformylglycinamide formyltransferase 2 {Bacillus subtilis}43.365.11361114633phosphoribosylglycinamide formyltransferase 2 {Bacillus subtilis}61.875.9155580156431ribose-phosphate pyrophokinase {Haemophilus influenzae}34.155.5	MJ0616	925486	925941	phosphoribosylaminoimidazole carboxylase (Methanobrevihacter emithil)	6 9 3	55	130	
1277597 1276734 phosphoribosylformylglycinamidine cyclo-ligase (Bacillus subtilis) 51.0 69.1 1507541 1507071 phosphoribosylformylglycinamidine synthase I (Bacillus subtilis) 52.9 71.5 262585 264714 phosphoribosylformylglycinamidine synthase II (Bacillus subtilis) 43.3 65.1 13611 14633 phosphoribosylglycinamide formyltransferase 2 (Bacillus subtilis) 61.8 75.9 155580 156431 ribose-phosphate pyrophosphokinase (Haemophilus influenzae) 34.1 55.5	MJ1592	1572482	1572009	Thorntoning in the state of the	20.3	7.0/	400	:
12775971276734phosphoribosylformylglycinamidine cyclo-ligase {Bacillus subtilis}42.764.415075411507071phosphoribosylformylglycinamidine synthase I {Bacillus subtilis}52.971.5262585264714phosphoribosylformylglycinamidine synthase II {Bacillus subtilis}43.365.11361114633phosphoribosylglycinamide formyltransferase 2 {Bacillus subtilis}61.875.9155580156431ribose-phosphate pyrophokinase {Haemophilus influenzae}34.155.5				Processor annihilazoresuccinocarboxamide synthase (Bacillus subtilis)		69.1	474	-41
15075411507071phosphoribosylformylglycinamidine synthase I {Bacillus subtilis}52.971.5262585264714phosphoribosylformylglycinamidine synthase II {Bacillus subtilis}43.365.11361114633phosphoribosylglycinamide formyltransferase 2 {Bacillus subtilis}61.875.9155580156431ribose-phosphate pyrophokinase {Haemophilus influenzae}34.155.6	MJ0203	1277597	1276734	phosphoribosylformylglycinamidine cyclo-ligase (Bacillus subtilis)	Γ	64.4	0,674	<u> </u>
262585 264714 phosphoribosylformylglycinamidine synthase II (Bacillus subtilis) 43.3 65.1 13611 14633 phosphoribosylglycinamide formyltransferase 2 (Bacillus subtilis) 61.8 75.9 155580 156431 ribose-phosphate pyrophosphokinase (Haemophilus influenzae) 34.1 55.5	MJ1648	1507541	1507051	phosphoribosylformylglycinamidine synthase I (Bacillus embrilie)	1		÷00	
13611 14633 phosphoribosylglycinamide formyltransferase 2 (Bacillus subtilis) 43.3 65.1 155580 156431 ribose-phosphate pyrophosphokinase (Haemophilus influenzae) 34.1 55.5	MJ1264	262585	264714	phosphoribosylformylglycinamidine synthase II (Bacillian Contain)	十	C.1.	4/1	
155580 156431 ribose-phosphate pyrophosphokinase {Haemophilus influenzae} 61.8 75.9	MJ1486	13611	14613	Though it begins subtility	7	65.1	2130	
155580 156431 ribose-phosphate pyrophosphokinase (Haemophilus influenzae) 34 1 55 5			655	prospiroringsylgiyeinamide formyltransferase 2 (Bacillus subtilis)	8.19	75.9	1023	
	MJI 366	155580	156431	ribose-phosphate pyrophosphokinase (Haemophilus influenzae)	34.1	\$ \$ \$	658	

Pyrimidine ri	bonucleotid	Pyrimidine ribonucleotide biosynthesis					_
MJ1581	1581578	1580661	aspartate carbamovitransferace cataloxic chain (Ecchanichia				
MIIAOC	973701			50.0	70.7	816	
0041 CM	104548	104183	aspartate carbamoyltransferase regulatory chain (Escherichia coli)	101	1 39	326	Τ-
MJ1378	145461	144037	carbamoyi-phosphate synthase, large chain (Bacillus cubrilia)		3	200	
M11381	143003	300.7	(Carrier Sancial)	29.7	80.0	1425	
1061614	143097	141328	carbamoyl-phosphate synthase, pyrimidine-specific, large subunit {Bacillus caldolyticus}	54.7	75.7	1770	
M1019	523003	522041					_
	223003	322041	carbamoyl-phosphate synthase, small chain (Bacillus subtilis)	49.6	1 69	190	_
MJ1174	358774	360279		663			
MJ0656	888785	202080		20.7	7.t.C	1506	
	6000	000000	cylidylate kinase (Bacillus subtilis)	310	2 03	400	
MJ1490	8032	6764	dihydroorotase {Bacillus caldolyticus}	1	5.5	480	
MIDEEA	0,,000	10000		34.5	56.3	1269	-9
45000M	289447	890284	dihydroorotase dehydrogenase (Bacillus subtilis)	43.1	7 7 7 7	643	<u>დ-</u>
MJ0293	1196756	9619611	thymidylate kinase (Schizosaccharomyces nombol	†	23	6	
90				31.2	58.7	561	
MJ1109	421875	421348	uridine 5'-monophosphate synthase {Dictyostelium discoldeum}	18.7	13	600	, -
MJ1259	271220	270543		7	0.4.0	328	
	2112	210743	uridylate kinase {Haemophilus influenzae}	77.6	-		

Salvage of nucleosides and nucleotides	cleosides an	d nucleotides				
MJ1459	43987	42413	adenine deaminase (Bacillus cubtilia)			
MJ1655	1499440	1499075	odenine monthe.it	35.9	61.7	1575
MJ0060	1412894	1412130	definite prospiretions (Hacmophilus influenzae)	35.8	62.5	366
M10667	870550	621017	metnyttnioadenosine phosphorylase {Homo sapiens}	41.3	63.2	756
8	0006/0	0/8150	thymidine phosphorylase (Mycoplasma genitalium)	30.5	52.2	1401
Sugar-nucleotide biosynthesis and conversion	ride biosynti	nesis and con	versions			
MJ1101	430386	429235	glucose-1-phosphate thumidala language			
MJ1334	188314	180084	TIDE 1.	32.0	56.0	1152
Repulatory functions	netions		Our-Blucose pyrophosphorylase (Mycoplasma genitalium)	42.7	63.6	171
7410800						
000000	/48410	747352	activator of (R)-2-hydroxyglutaryl-CoA dehydratase {Acidaminococcus	31.8	51.2	1059
MJ0004	1466944	1466288				
		66200	activator of (K)-2-hydroxyglutaryl-CoA dehydratase (Acidaminococcus fermentans)	39.0	61.1	069
MJ1344	180975	181229	nitrogen regulatory protein P.11 / Hasma-Lilas			
MJ0059	1413301	1413047	nitrogen een lee	56.5	73.0	255
MJ0300	1188833	10011	gc.i icguiatory protein P-II {Haemophilus influenzae}	56.5	75.3	255
	7000011	1100194	putative transcriptional regulator (Bacillus subtilis)	27.8	503	630
MJ0151	1325766	1325323	putative transcriptional regulator (Pyrococcus furiosus)			ŝ.
MJ0723	815573	815190	Dutative transcriptional seculators (P.	2.1.0	02.0	444
			remaind the state of the state	¢13	92.2	30.4

			Replication			
Degradation of DNA	of DNA					
MJ1434	68536	68048	endonuclease III (Bacillus subtilis)	78.7	1 85	760
MJ0613	927393	928424	endonuclease III {Bacillus subtilis}	1 2	, ,	1601
MJ1439	65786	65208	thermonuclease precursor {Staphylococcus hyicus}	36.0	5	7501
DNA replication, restriction, modification, r	ion, restricti	lon, modificat	ion, recombination, and renair	20:0	7.	6/6
M11020	\$10622	20003				
10101	210033	209812	dimethyladenosine transferase {Bacillus subtilis}	38.4	58.8	759
MJ0104	1373055	1371130	DNA helicase, putative (Homo sapiens)	35.2	26.7	1076
MJ0171	1297428	1299053	DNA ligase (Desulfurolobus ambivalens)	35.8	62.4	1636
MJ0869	680404	679445	DNA repair protein {Saccharomyces cerevisiae}	44.6	600	070
MJ1444	58945	58052	DNA repair protein RAD2 {Homo sapiens}	37.3	2.20	200
MJ0254	1232179	1231757	DNA renair protein BADS1	5/,5	03.3	894
77,002,1	202023		City tepail protein (ADD) (Homo sapiens)	32.5	58.4	423
10600141	2/9280	5//424	DNA replication initiator protein (Xenopus laevis)	28.1	40.0	2157
MJ1652	1503610	1501559	DNA topoisomerase I (Mycoplasma genitalium)	34.0	55.0	2052
MJ0885	656470	096099	DNA-dependent DNA polymerase family B (Pyrococcus sp.)	47.3	0 8 9	4401
MJ1529	1630880	1630413	methylated DNA protein cysteine methyltransferase (Haemophilus influenzae)	35.0	2,79	469
MJ1498	1548	715	modification methylase (Haemophilus parainfluenzae)	3.5	;	100
MJ0598	942522	941860	modification methylase (Haemophilus influenzae)	3. 4	53.8	750
MJ1328	193775	186761	modification methylase (Haemophilus influenzae)	1	1 3	66
				_	-	\o\

	٥	7070		r	T			Τ-	_			_	_		<u>- 93</u>	<u>5-</u>	;					PCI	/US	97/1	1490
		789	1035	852		507	1887	624		627	5424	1074		1683	540	699		4839	639	1673	3477	1	CHS	3125	645
		56.2	56.7	73.0		63.9	54.7	54.3		99.	64.6	49.2	3	24.4	61.1	59.2];	97.4	64.6	61.4	52.2	333	7,,,,	59.9	57.9
		34.7	39.7	54.5	١	40.3	31.9	31.5	;	48.1	45.2	32.5	320	247	37.3	29.3	41.0	0.7	45.2	39.7	31.1	20,5		36.2	38.4
	modification methylase (Methanshastarium 1	modification methylace (Theories and Theories and Theorie	modification (1) (1) (2) (2) (2) (3) (3) (4) (4) (4) (4) (4) (4) (4) (4) (4) (4		mutator mutT protein {Escherichia coli}	probable ATP-dependent helicase (Unamerica)	nroliferating self	Frontianing-cell nuclear antigen (Saccharomyces cerevisiae)	Proliferating-cell nucleolar antigen, 120 kDa (Homo saniens)	replication factor C (Homo saniens)	replication factor Classes	(Homo sapiens)	restriction modification enzyme, subunit M1 (Mycoplasma pulmonis)	restriction modification enzyme suburit Act (a.c.	restriction modification		reverse gyrase (Sulfolobus acidocaldarius)	ribonuclease HII (rnhB) (Escherichia colin	type I restriction envires ECOB 13475 000	type I restriction and the control of the control o	The result of the season of th	type I restriction enzyme (Haemophilus influenzae)	type I restriction enzyme (Haemonhilus influenzae)	type I restriction and managed in the contract of the contract	surction enzyme Cirl, specificity subunit (Citrobacter freundii)
	975309	327248	555896	384346	384248	598916	1237322	1446324	477C+L1	84727	696099	310102	210102	1345548	1347179	1640747	75/0501	1341939	54271	1352847	215020	979616	49456	1628493	
	974521	326214	555045	181747	74/506	600802	1237945	1444508		79304	662042	308420		1345009	1346511	1653580	20000	1341301	55944	1349371	213714	11.51.5	52581	1629137	
7,105.73	MJ0563	MJ1200	MJ0985	MJ1149		MJ0942	MJ0247	MJ0026		MJ1422	MJ0884	MJ1220	Miniss	M20132	MJ0130	MJ1512		MJ0135	MJECL42	MJ0124	MJ1214	27.10.17	MJECL40	MJ1531	

MJ1218	310547	311776	type I restriction-modification enzyme, S subunit {Escherichia coli}	70.7	40.7	9551
MJ0984	556397	555909	type II restriction enzyme (Methanobacterium thermoformicioum)	7.5.7	47.7	0621
MJ0600	940932	940315	type II rectriction on the David Co.	£3.9	7./0	489
			ype it issuiction enzyme Drivii (Streptococcus pneumoniae)	46.0	67.4	618
			Transcription			
DNA-dependent RNA polymerases	nt RNA pol	lymerases				
MJ1042	407715	401737				
		475/52	UNA-dependent RNA polymerase, subunit A' {Methanococcus vannielii}	74.5	88.1	3984
MJ1043	493546	491078	DNA-dependent RNA polymerase, subunit A" {Methanococcus vannielii}	66.7	83.5	2460
MJ1041	499305	497866	DNA-dependent RNA polymerase, subunit B' {Methanococcus vannielii}	76.3	2 6	246
MJ1040	501124	499862	DNA-dependent RNA polymerase, subunit B" (Methanococcus vannielii)	1 2	2 2	0440
MJ0192	1283621	1283148	DNA-dependent RNA polymerase, subunit D (Arabidoneie thaliana)	7.5.7	4.70	507
MJ0397	1113901	1114371	DNA-denendent RNA notumeror	c.kr	38.6	474
M10306			serior of the se	47.9	8.02	471
0450LM	1114384	1114560	DNA-dependent RNA polymerase, subunit E" (Sulfolobus acidocaldarius)	35.9	603	133
MJ1039	501599	501366	DNA-dependent RNA polymerase, subunit H (Methanococcus vannielii)	707	;	, , ,
MJ1390	134111	134350	DNA-dependent RNA polymerase, subunit 1 (Sulfolohus aridocaldarius)	1	3	234
MJ0197	1281417	1281247	DNA-dependent RNA polymerase subunit K (Haloaroula mariamas)	+	6.5	240
MJ0387	1119216	1119512	DNA dependent BNA soluments	45.5	83.3	171
201012			Surforces (Surforces) Subunit (Surfolobus acidocaldarius)	35.6	63.4	297
M30196	1281779	1281561	DNA-dependent RNA polymerase, subunit N (Haloarcula marismortui)	53.8	83.4	210

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Transcription factors	n factors						
MJ0941	601867	600923	Dufative francoriusion : ::::				
MJ1045	490363	489848	mitative transcription initiation factor IIIC (Saccharomyces cerevisiae)	20.1	44.1	945	Γ
			vannielii)	47.9	73.7	516	
MJ0372	1134509	1134123	putative transcription termination and it				
MJ0507	1024170	1024631	TATA-binding transcription in it.	38.6	63.8	387	
MJ0782	766586	768592	transcription initiality and actor { Thermococcus celer}	51.4	74.0	462	
MJ1148	384277	384567	removable initiation factor IIB (Pyrococcus woesei)	63.8	77.6	2007	T
RNA processing	Bu		"anscription-associated protein, ('TFIIS') {Thermococcus celer}	56.4	69.0	291	7
MJ0697	849814	849125	fibrillerin-like				
Transfation			Methanococcus vannielii}	75.3	88.3	069	T-
MJ0160	1308036	1309265	PET112 mendein (C				,
Amino acyl tRNA synthetases	NA synthet	ases	(Saccharomyces cerevisiae)	32.3	53.7	1230	7/
MJ0564	971657	974149	alancist NA ******			:	_
MJ0237	1244137	1242641	aroinvi (BNA cont.	28.0	53.1	2493	_
MJ1555	1605935	1604679	Benary (10NA)	31.3	52.7	1497	_
MJ1377	145796	147325	chitemal above	87.8	75.6	1257	,
MJ0228	1253254	1251524	glycyl-tRNA synthetiase (Methanobacterium thermoautotrophicum)	51.7	73.6	1530	
MJ1000	543634	542396	hieride DNA section (300)	45.8	65.2	1731	·
		7	mistray - traday synthetase (Streptococcus equisimilis)				

MJ0947	591914	594817	isoleucyl-tRNA synthetase (Methanobacterium thermoautotrophicum)	52.1	70.0	2004
MJ0633	912642	910015	leucyl-tRNA synthetase (Saccharomyces cerevisiae)	34.4	840	8696
MJ1263	266697	264745	methionyl-tRNA synthetase {Haemophilus influenzae}	35.6	86.0	1053
MJ0487	1041343	1039994	phenylalanyl-tRNA synthetase, subunit alpha {Saccharomyces cerevisiae}	919	5 2	1360
MJ1108	423555	425198	phenylalanyl-tRNA synthetase, subunit beta {Saccharomyces cerevisiae}	3 2	2.5	1530
MJ1238	287985	289172	prolyf-tRNA synthetase {Homo sapiens}	30.5	\$05	1044
MJ1197	332116	330257	threonyl-tRNA synthetase {Synechocystis sp.}	20.5	5.50	0071
MJ1415	96418	95369	tryptophanyl-tRNA synthetase (Schizosaccharomyces pombe)	30.5	55.1	1000
MJ0389	1118380	1117616	tyrosyl-tRNA synthetase (Homo sapiens)	30.0	63.7	768
MJ1007	536642	534186	valyl-tRNA synthetase {Bacillus stearothermophilus}	36.1	2,75	2463
Degradation of proteins, peptides, and glycopeptides	f proteins, p	eptides, and	glycopeptides		200	
MJ1176	356300	357370	ATP-dependent 26S protease regulatory subunit 4 (Homo canions)	6:3		8-
M11494	4302	\$177	ATP	21.0	4.1	1/21
	7000	2123	A 17-dependent 205 protease regulatory subunit 8 (Methanobacterium thermoautotrophicum)	58.6	78.2	822
MJ1417	93716	91932	ATP-dependent protease La (Bacillus brevis)	3.7 g	623	1785
MJ0090	1387867	1386755	collagenase (Porphyromonas gingivalis)	3.6	3	113
MJ1130	400455	401969	O-sialoglycoprotein endopeptidase {Saccharomyces cerevisiae}	20.5	27.0	515
MJ0651	891988	892842	protease IV {Haemophilus influenzae}	35.0	6,19	CICI
MJ0591	947601	946861	proteasome, subunit alpha {Methanosarcina thermorbila}	2.50	7.00	650
				ر: ادرات	0.0	141

289440 2	289967	proteasome, subunit beta (Methanosarcina thermonkila)				
742381 7	743364	Xaa-pro dipeptidase (Lactobacillus delbusatii)	47.5	68.2	528	
547987	546635	Zn professe (Haemonkillus in Discourse)	36.1	65.2	984	
1			33.9	55.0	1353	
733804	734793	deoxyhypusine synthase (Home social)				
253925	254653	diphthine synthage (Services aprens)	20.0	70.7	.990	_
1296723	1297175	1 isosement	40.7	61.5	729	_
192979	192098	methionine aminocarii ta care (Escherichia coli)	47.6	59.4	453	
1630123	1629764	N-terminal aceta-ti	36.2	55.1	882	
ו ה	1573833 1573072	selenium donos	39.7	55.7	360	
	≝ ⊦	diffication	34.3	57.1	762	
ı vo	1022576 1023502	المارات				
1240163	┽╾	acture itoosomal protein P0 (L10E) {Methanococcus vannielii}	63.2	82.1	726	
. .	0770171	ribosomal protein HG12 (Catus (cat))	63.7	81.9	99	_
- 1	325460	ribosomal protein HS6-type (Haloarcula marismortui)	47.0		36.	
1021912	1022460	ribosomal protein L1 {Methanococcus vannieliis	2: ;	5	100	
1133926	1133540	Tibosomal protein [11] Sc. (61)	64.5	80.3	549	
1 -	20000		47.2	72.4	387	
1023032	1023937	ribosomal protein L12 (Methanococcus vannielii)	72.8	80.0	306	
1282568	1282260	ribosomal protein L13 (Haloarcula marismortui)				
1058694	1058452	ribosomal protein L14 (Methanococus varnicatis)	44.9	66.4	309	
		Control of the contro	78.8	97.5	243	

MJ0657	888216	887977	ribosomal protein L14B (Saccharomyces cerevisiae)			
MJ0477	1052625	1052302	ribosomal protein L.15 (Methanococcus manifelia)	36.4	29.8	240
MJ0983	556982	557290	ribosomal protein 1.158 (Thermonlocation 1.1.1.	62.7	79.5	324
MJ0474	1054523	1053939	ribosomal protein 1 18 (Marks	62.3	78.6	309
MJ0473	1054978	1054559	ribosomal account to the control of	73.3	84.3	585
MJ0179	1291786	1291052	ribosomal profess 1.3 (Methanococcus vannielii)	67.0	86.4	420
MJ0040	1431958	1432260	ribosomal activity of the state	74.0	87.0	735
MJ0460	1061493	1061080	ileania piotein LZ I (fraioarcula marismortui)	54.5	62.3	303
M10170		6001001	ribosomai protein L22 (Haloarcula marismortui)	40.7	61.7	405
8/100M	1292097	1291840	ribosomal protein L23 (Methanococcus vannielii)	8.69	919	258
M1040/	1058340	1058062	ribosomal protein L24 {Methanococcus vannielii}	20.5	93.0	
MJ1201	325929	326078	ribosomal protein L24E (Haloarcula marismortui)	6.07	0.50	617
MJ0462	1060388	1060212	ribosomal protein 1.29 (Halohanterium halatina)	54.6	66.7	150
MJ0193	1283076	1282705	ribosomal protein 1 30E 11.1	51.0	6.69	177
MJ0176	1293794	1202014	(ia	48.7	68.7	372
MJ1044	490704	490399		45.2	63.9	198
MJ0049	1421907	1422152	nelii)	63.9	84.1	306
MJ0472	1055464	1055063		40.9	66.2	246
MJ0655	889197	888931	ribosomal protein 134 (Action 1988)	58.0	77.4	402
MJ0098	1380525	1380686		36.8	58.3	267
		7	ital protein L3 / {Letshmania infantum,}	50.0	67.4	791

MJ0593	945958	945683	in the state of th				
MJ0177	1292889	+	ribosomal protein L3/8 (Homo sapiens)	44.6	58.7	276	
MJ0707	838122	838229	ribosomal protein 140 (Co. 1	49.4	66.3	756	
MJ0249	1236729	1236448	ribosomal protein 144 (11-1-1-1-1-1-1-1-1-1-1-1-1-1-1-1-1-1-1	57.6	66.7	108	
MJ0689	854995	855150	ribosomal protein 146 (Haloarcula marismortui)	38.8	58.1	282	
MJ0469	1057259	+	ribocomal protein 1.6 (M. 4)	52.0	70.0	156	
MJ0471	1056071	1055526	ribocomal profession 17 (Methanococcus vannielii)	72.5	84.5	537	
MJ0476	1053137	┼─	ribosomal protein L9 (Methanococcus vannielii)	66.5	82.5	546	
MJ0595	944670	944473	ribosomal protein I V 18-16-1-1	70.3	88.6	393	
MJ0322	1172916	1173218	ribosomal protein C10 to.	38.9	66.7	198	
MJ0191	1283956	1283735	ribosome profession of trytococcus woeses	67.0	91.0	303	-/
MJ1046	489559	489260	ribosomal protein S11 {Haloarcula marismortui}	67.2	80.0	222	0/-
MJ0036	1434801	1434352	ribosomal protein 512 (Methanococcus vannielii)	87.0	96.0	300	
MJ1474	26554	26054	ribosomal protein S13 (Brugia pahangi,)	49.4	71.0	450	
MJ0465	1059233	1058883	ribosomal protein S17 (Master	21.7	48.2	501	
MJ0245	1238750	1238896	ribosomal profein S17B (Seconds Vannielii)	71.6	82.4	351	
MJ0189	1285220	1284771	al Drotein S18 (Arabidonei: 41-11-11-11-11-11-11-11-11-11-11-11-11-1	55.4	80.9	147	PC
MJ0180	1290861	1290508	:	42.3	68.5	450	1705
MJ0692	853669	854046		56.9	73.3	354	97/14
			STATE				\$ 5

MJ0394	1115064	1115366	ribosomal protein S24 (Haloarcula marismortui)	42.6	64 4	203
MJ0250	1236377	1236192	ribosomal protein S27 {Saccharomyces cerevisiae}	42.6	62.9	201
MJ0393	1115369	1115548	ribosomal protein S27A {Caenorhabditis elegans}	58.4	0.55	100
MJ0461	1061060	1060437	ribosomal protein S3 (Haloarcula marismortui)	60 1	22.1	234
MJ1202	325575	325808	ribosomal protein S33 {Kluyveromyces lactis}	2 5	1 1 1 8	73.4
MJ0980	1928361	559252	ribosomal protein S3a {Catharanthus roseus}	20.8	2 2	407
MJ0190	1284710	1284150	ribosomal protein S4 {Sulfolobus acidocaldarius}	513	7 89	764
MJ0468	1057935	1057318	ribosomal protein S4E (Methanococcus vannielii)	70.0	24.5	100
MJ0475	1053877	1053275	ribosomal protein S5 {Methanococcus vannielii}	75.7	200	810
MJ1260	270075	269683	ribosomal protein S6 {Homo sapiens}	25.	0.00	
MJ0620	922671	921799	ribosomal protein S6 modification according (1122)	7.00	28.0	565
MILOOI	64222	-07.173		34.4	57.3	873
1001011	177740	24148/	ribosomal protein S6 modification protein II (Haemophilus influenzae)	24.8	47.4	741
MJ1047	489046	488627	ribosomal protein S7 (Methanococcus vannielii)	65.8	83.6	420
MJ0470	1056445	1056113	ribosomal protein S8 {Methanococcus vannielii}	71.7	60,0	3 :
MJ0673	873106	872720	ribosomal protein S8E {Haloarcula marismortui}	1	7.60	252
MJ0195	1282118	1281840	ribosomal protein S9 {Haloarcula marismortui}	十	75.0	38/

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| tRNA modification   | cation  | -       |                                                                                 |      |       |          | F           |
|---------------------|---------|---------|---------------------------------------------------------------------------------|------|-------|----------|-------------|
| MJ0946              | 595006  | 596040  | N2.N2-dimethylonanosine (BNA moth.)                                             |      |       |          |             |
| MJ1675              | 1478684 | 1477755 | nsendouridulate constant 1 (1)                                                  | 31.6 | 56.0  | 1035     |             |
| MJ0436              | 1081116 | +-      | recently rate symmate 1 (Tracmophilus influenzae)                               | 33.5 | 57.2  | 930      |             |
| Translation factors | Betors  | 1002132 | queuine trans ribosyltransferase {Escherichia coli}                             | 30.4 | 47.6  | 1617     |             |
| MJ0829              | 773534  | 2227.0  |                                                                                 |      |       |          | _           |
|                     | PCCC2/  | 09777/  | peptide chain release factor, eRF, subunit 1 (Xenopus laevis)                   | 33.0 | 573   | 1336     | <b>T</b>    |
| MJI 505             | 1659133 | 1661085 | putative ATP-dependent RNA helicase, eIF-4A family {Saccharomyces               | 30.8 | 51.9  | 1953     | 1           |
| MJ1574              | 1587062 | 1588927 | nitative ATD desired                                                            |      |       |          |             |
| MJ0669              | 876636  | 877617  | printed Att - dependent KNA helicase, elf-4A family (Bacillus subtilis)         | 33.1 | 56.0  | 1866     |             |
| M10405              | 1036437 |         | putative A 1 F-dependent RNA helicase, elF-4A family (Bacillus subtilis)        | 44.5 | 65.8  | 1002     | _           |
|                     | 1033432 | 1034044 | putative translation factor, EF-TU/1 alpha family (Thermus aquaticus)           | 36.0 | 0 33  | 965      | _           |
| MJ0262              | 1225060 | 1221653 | putative translation initiation factor, FUN12/bIF-2 family (Saccharomyces       | 39.3 | 61.5  | 3408     | <u>-/03</u> |
| MJ0324              | 1171774 | 1177820 |                                                                                 |      |       | <b>?</b> |             |
|                     | 57/1/11 | 11/2830 | translation elongation factor, EF-1 alpha (Methanococcus vannielii)             | 9 9  | 9 9 9 |          | _           |
| MJ1048              | 488471  | 486336  | translation elongation factor FF-2 / Maihanne                                   | /8.9 | 90.8  | 1107     |             |
| MJ0445              | 1073262 | 1073483 | translation initiation factor alt 14 th                                         | 74.8 | 88.5  | 2136     |             |
| MJ0117              | 1357516 | 1358196 | franclation initiation features in a sucremoplasma acidophilum)                 | 52.8 | 70.3  | 222      |             |
| MJ0097              | 1380885 | 1381313 | francisco initiation factor, elf-2, subunit alpha (Saccharomyces cerevisiae)    | 32.2 | 56.5  | 189      |             |
| MJ1261              | +-      | 269164  | ransiation initiation factor, eIF-2, subunit beta {Drosophila melanogaster}     | 32.1 | 60.4  | 429      |             |
| MIDACA              | +       | 50007   | translation initiation factor, eIF-2, subunit gamma {Homo sapiens}              | 52.6 | 71.9  | 1233     |             |
| *C*****             | 1066217 | 1067065 | translation initiation factor, eIF-2B, subunit alpha (Saccharomyces cerevisiae) | 37.9 | 56.4  | 840      |             |
|                     |         |         |                                                                                 | 1    |       | 22,      | _           |

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| MJ0122                         | 1353264      | 1354127 | translation initiation factor, elF-2B, subunit delta {Mus musculus}     | 20.4 | 3 7 7    | 770  |
|--------------------------------|--------------|---------|-------------------------------------------------------------------------|------|----------|------|
| MJ1228                         | 300895       | 301236  | translation initiation factor, eIF-5a (Sulfolobus acidocaldarius)       | 5.62 | 5 5      | 100  |
| Transport and binding proteins | binding pr   | oteins  |                                                                         | 20.0 | 09.7     | 342  |
| MJ0719                         | 818577       | 820289  | ABC transporter ATP hinding access:                                     |      |          |      |
| 111033                         | 1 250        |         |                                                                         | 49.6 | 6.99     | 1713 |
| M31023                         | 218606       | 517821  | ABC transporter ATP-binding protein (Bacillus firmus)                   | 49.2 | 72.4     | 786  |
| MJ1572                         | 1590114      | 1589518 | ABC transporter ATP-binding protein {Mycoplasma genitalium}             | 000  | 97.6     |      |
| MJ0035                         | 1435236      | 1435829 | ABC transporter subunit {Cyanelle Cyanophora}                           | 2.5  | c:/8     | / k  |
| MJ1508                         | 1656015      | 1655446 | ABC transmission and Link ATR 1: 1:                                     | 33.9 | <u>~</u> | 294  |
|                                |              |         |                                                                         | 45.7 | 68.3     | 570  |
| MJ1332                         | 189987       | 191117  | GTP-binding protein (Saccharomyces cerevisiae)                          | 38.7 | \$ 0 y   | 1131 |
| MJ1326                         | 196392       | 195292  | GTP-binding protein {Schizosaccharomyces pombe}                         |      | 2        | 1611 |
| MJ1408                         | 103440       | 102430  |                                                                         | 4:15 | ?        | 101  |
|                                | , Land       | 102430  | O 17-binding protein, G 1 P 1/OBG-family (Saccharomyces cerevisiae)     | 30.5 | 58.4     | 1020 |
| MJ1464                         | 39865        | 38858   | hypothetical GTP-binding protein (SP:P40010) {Saccharomyces cerevisiae} | 33.0 | Ĭ        | 1000 |
| MJ1033                         | 5072:74      | 506324  | magnesium and cobalt transport protein (Haemonhilus influence)          | 2 2  |          | 9001 |
| MJ0091                         | 1386551      | 1385751 | Na+/Ca+ exchanger protein (Escherichia coli)                            | 42.2 | 6.7.9    | 951  |
| M10282                         | $\mathbf{T}$ | 1303673 |                                                                         | 32.3 | 58.6     | 801  |
| (4)70702                       | 1204330      | 1203563 | nucleotide-binding protein (Homo sapiens)                               | 47.5 | 0 89     | 876  |

| Amino acids, peptides and amines | peptides an | d amines |                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                |       |       |      |
|----------------------------------|-------------|----------|--------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|-------|-------|------|
| MJ0609                           | 933328      | 934587   | amino acid transporter {Arabidopsis thaliana}                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  | 3     |       |      |
| MJ1343                           | 181359      | 182519   | ammonium transport protein AMT1 (Arahidonsis shaliana)                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         | 21.9  | 48.7  | 1260 |
| MJ0058                           | 1413598     | 1414770  | ammonium transporter (Escherichia coli)                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        | 33.6  | 23.3  | 1161 |
| MJ1269                           | 258901      | 257993   | branched-chain amino acid transnort protein light feathering                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   | 34.2  | 52.2  | 1173 |
| MJ1266                           | 261404      | 260577   | hranched wheir amino and a second month (Escuential Coll)                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      | 30.8  | 24.6  | 606  |
| M11270                           | 267005      | 26,000   | orange chain annua acid transport protein liv1 {Escherichia coli}                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              | 28.8  | 55.2  | 828  |
|                                  | 060/67      | 230934   | branched-chain amino acid transport protein livM (Escherichia coli)                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            | 28.7  | 52.2  | 696  |
| MJ1196                           | 332430      | 333311   | cationic amino acid transporter MCAT-2 (Mus musculus)                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          | 246   | \$0.5 | 663  |
| MJ0304                           | 1185908     | 1186333  | ferripyochelin binding protein {Pseudomonas aeruginosa}                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        | 3 3   | 2,25  | 700  |
| MJ0796                           | 752786      | 752118   | Slutamine transport ATP-binding protein O (Escherichia colly                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   | 200   |       | 074  |
| MJ1267                           | 260465      | 259707   | high-affinity branched-chain amino acid transport ATD hinding                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  | 4/.9  | 27.0  |      |
|                                  |             |          | {Pseudomonas aeruginosa}                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       | 34.2  | 8.09  | 759  |
| MJ1268                           | 259458      | 258973   | high-affinity branched-chain amino acid transport ATP-binding protein (Salmonella typhimurium)                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 | 40.4  | 68.6  | 486  |
| Anions                           |             |          |                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                |       |       |      |
| MJ0412                           | 1099862     | 8090011  | nitrate transport ATP-binding protein {Synechococcus sn}                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       |       | 9     |      |
| MJ0413                           | 1099017     | 1099826  | nitrate transport permease protein {Synechococcus sn}                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          | 44.0  | , S   | /4/  |
| MJ1012                           | 529685      | 530431   | phosphate transport system ATP-binding protein (Escherichia colit                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              | 7.4.5 | 99.4  | /50  |
| MJ1013                           | 528941      | 529642   | phosphate transport system permease protein A (Haemonhilus in duant                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            | 99.9  | 80.7  | /47  |
| MJ1014                           | 528397      | 528810   | phosphate transport system nermeage protein ( the protein )                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    |       | 50.0  | 702  |
|                                  |             |          | (nacmopaline in the interest of the interest o | 40.0  | 66.5  | 414  |

| MJ1009 | 532458       | 533165                                     | the contract of the contract o |        |        |      |
|--------|--------------|--------------------------------------------|--------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|--------|--------|------|
| 1      |              | 20122                                      | phospitate transport system regulatory protein (Escherichia coli)                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              | 28.5   | 54.6   | 208  |
|        | 526871       | 527698                                     | phosphate-binding protein (Xanthomonas oryzae)                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 | 45.8   | 603    | 920  |
| rate   | s, organic a | Carbohydrates, organic alcohols, and acids | ıcids                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          |        | 7.00   | 070  |
|        | 960439       | 959399                                     | malic acid transport protein (Schizosaccharomycan amelic                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       |        |        |      |
|        | 786703       | 787524                                     | malic acid transport protein (Schizosachasomas | 23.8   | 47.9   | 1041 |
|        | 1354728      | 1355291                                    | SN-g Veerol-3-phoenhate transport ATD 1: 1:                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    | 26.5   | 49.3   | 822  |
|        | 206861       | 205926                                     | sodium denerdent to the control of t | 33.4   | 51.7   | 564  |
|        |              |                                            | Section acpellation in the description of the second philas influenzae)                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        | 37.8   | 61.0   | 936  |
|        | 444480       | 446333                                     |                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                |        | ļ      |      |
|        | 004444       | 443773                                     | cobalt transport ATP-binding protein O (Salmonella typhimurium)                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                | 46.1   | 9.99   | 744  |
| I      | 443372       | 443527                                     | cobalt transport protein N (Salmonella typhimurium)                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            | 59.1   | 70 6   |      |
| T      | 443778       | 444374                                     | cobalt transport protein Q (Salmonella typhimurium)                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            | 300    |        | -/06 |
|        | 1388820      | 1388059                                    | ferric enterobactin transport ATP-binding protein (Escherickia                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 | 7.6.9  | 25.6   | ļ    |
|        | 674824       | 674123                                     | ferric enterobactin transport A TP-hinding access (1771)                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       | 33.1   | 29.6   | 762  |
|        | 967842       | 0,40,867                                   |                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                | 31.5   | 60.3   | 702  |
|        |              | 10000                                      | ictions from transport protein B {Escherichia coli}                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            | 35.8   | 61.2   | 2016 |
| 1      | 0/0239       | 670442                                     | hemin permease (Haemophilus influenzae)                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        | 37.0   | 3      | 1    |
|        | 1390284      | 1389385                                    | hemin permease (Yersinia enterocolitica)                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       | $\top$ | +      | 204  |
|        | 1392668      | 1391613                                    | iron transport system binding profein (Becillie subvilie)                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      | $\top$ | $\top$ | 900  |
|        | 670677       | 671498                                     | iron(III) dicitrate transport evetem permanant                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 | 32.9   | 53.3   | 1056 |
|        | 64080        | 60403                                      | magnetic - Constitution of the control of the contr | 30.8   | 52.8   | 822  |
| 1      |              | 7,22                                       | inaginesium eneratase subunit (Arabidopsis thaliana)                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           | 35.3   | 57.3   | 3678 |

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| MJ0911 | 628932  | 629972  | magnesium-chelatace cubumis (E)                                  |      |      |      |         |
|--------|---------|---------|------------------------------------------------------------------|------|------|------|---------|
| MJ1275 | 253661  | 252597  | NA(+)/H(+) entirement (Eugliena gracilis)                        | 54.9 | 73.4 | 1041 |         |
| MJ0672 | 873748  | 874665  | Nat frances (11)                                                 | 29.8 | 59.9 | 1065 | $T^{-}$ |
| MJ1231 | 297233  | 298873  | oxaloacetate decet                                               | 39.3 | 63.1 | 918  | 1       |
| MJ1357 | 164247  | 165065  | Dutative notassium change and a subunit (Salmonella typhimurium) | 52.0 | 68.7 | 1641 | 7       |
| MJ1367 | 154669  | 155559  | Sulfate Dermease (CV&A) (C.mo.)                                  | 42.9 | 66.7 | 618  |         |
| MJ1368 | 153995  | 154666  | sulfate/thiosulfate transmit                                     | 38.5 | 64.5 | 168  | _       |
| MJ1485 | 16909   | 15713   | TRK system moteocium                                             | 30.9 | 59.4 | 672  | _       |
| MJ1105 | 426702  | 427217  | TRK system moteonium updake protein {Escherichia coli}           | 29.5 | 58.5 | 1197 |         |
| Other  |         |         | Journ Poussium uptake protein A (Methanosarcina mazei)           | 39.3 | 57.6 | 516  |         |
| MJ1142 | 390844  | 389885  | arenira numa dii                                                 |      |      |      | 70      |
| MJ0822 | 727897  | 729522  | ATPase, vanadate sensitive (Machine 1918)                        | 34.7 | 55.9 | 096  |         |
| MJ0718 | 820399  | 821523  | chromate recistance                                              | 48.1 | 0.69 | 1626 |         |
| MJ1226 | 304219  | 301988  | H+-transporting A TPase / Archidents (1.1.                       | 27.9 | 52.4 | 1125 |         |
| MJ1560 | 1600958 | 1601974 |                                                                  | 45.1 | 63.7 | 2232 |         |
|        |         |         | Staphylococcus aureus)                                           | 28.8 | 51.1 | 1017 |         |

| Other categories                      | ies           |              |                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                |      |       |      | Γ             |
|---------------------------------------|---------------|--------------|--------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|------|-------|------|---------------|
| MJ1365                                | 157333        | 156458       | pheromone shutdown protein (Enterococcus faecalis)                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             | 1    |       |      | 1             |
| MJECL24                               | 28069         | 28845        | SOI motein (Bonilles Little)                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   | 31.2 | 27.7  | 8/6  | Т             |
|                                       |               |              |                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                | 34.0 | 62.1  | 276  |               |
| Ding and analog sensitivity           | og sensitivi  | ć.           |                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                |      |       |      | Γ             |
| MJ1538                                | 1621434       | 1650691      | K. lactis toxin sensitivity protein KTI12 (Saccharomyces cerevisiae)                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           | 38.7 | 90,   |      | $\top$        |
| MJ0102                                | 1375563       | 1375859      | phenylacrylic acid decarboxylase (Saccharomyces cerevisies)                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    | 5 3  | 9.0   | /44  | $\top$        |
| Phage-related functions and prophages | functions a   | nd prophages |                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                | 20.0 | /4.0  | 767  | T             |
| MJ0630                                | 915023        | 914598       | Sodium-dependent phosphate francoorer (Crisselles                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              |      |       |      | 一             |
| Transnoson-rolated Grant              | loted Grant   |              | Suscina guscina                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                | 32.6 | 8.09  | 426  |               |
| - Pocodomi-                           | ומוכח וחווכוו | Ons          |                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                |      |       |      |               |
| MJ0367                                | 1138754       | 1138080      | integrase (Weeksella zoohelcum)                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                | 30 0 | 17.73 | 636  | $\overline{}$ |
| MJ0017                                | 1455555       | 1454946      | transposase {Bacillus thuringiensis}                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           | 200. | ş.    | 6/2  | 7             |
| Other                                 |               |              |                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                | 29.5 | 55.0  | 610  | 10            |
|                                       |               |              |                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                |      |       |      | 8-            |
| MJ1064                                | 466505        | 467095       | acetyltransferase {Escherichia coli}                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           | 47.0 | 3     |      | _             |
| MJ1612                                | 1549430       | 1548297      | BcpC phosphonopyruvate decarboxylase (Streptomyces hyarosconicus)                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              | 2,1  | 4.20  | 160  | <del></del>   |
| MJ0677                                | 868213        | 869160       | ethylene-inducible protein homolog (Heves brasiliensis)                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        | 31.1 | 48.9  | 1134 |               |
| MJ0534                                | 1003199       | 1002072      | flavonrotein (Methanohacterium themselven)                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     | 08.3 | 81.0  | 948  | -             |
| M10748                                | 707507        | 7007         | Commission of the management of the module o | 34.6 | 57.2  | 1128 |               |
|                                       | 131304        | /980/3       | Ilavoprotein {Methanobacterium thermoautotrophicum}                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            | 67.0 | 82.6  | 1170 | r             |
| MJ0256                                | 1230191       | 1229760      | fom2 phosphonopyruvate decarboxylase {Streptomyces wedmorensis}                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                | 36.7 | 58.5  | 432  | _             |
| MJ1682                                | 1472535       | 1473320      | heat shock protein X {Haemophilus influenzae}                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  | 30.4 |       | 2 2  | <del></del> - |

| MJ0866       | 682753  | 682367. | HIT protein, member of the HIT-family (Sarcharomycae caracicies)                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               | , 65   |      |       |             |
|--------------|---------|---------|--------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|--------|------|-------|-------------|
| MJ0294       | 1193529 | 1195817 | large helicase related protein. LHR (Escherichia coli)                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         | 39.4   | 84.8 | 387   | T-          |
| MJ0010       | 1460660 | 1459497 | phosphonopyruyate decarboxylase / Streeton                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     | 31.4   | 53.6 | 2289  |             |
| MJ0734       | 805855  | 806439  | The archein (Close 12:-                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        | 28.0   | 47.2 | 1164  |             |
| MJ0559       | 978787  | 077700  | constraint (Clostrigum perfringens)                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            | 48.9   | 69.2 | 585   | _           |
| M11100       | 431764  | 730,480 | sure survival protein {Escherichia coli}                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       | 34.7   | 55.6 | 862   |             |
| M10543       | 20,000  | 450489  | urease operon protein (Mycobacterium leprae)                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   | 33.2   | 55.0 | 1266  | _           |
| CHOOSE       | 79008/  | 991100  | Wilm's tumor suppressor homolog (Arabidopsis thaliana)                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         | 45.6   | 640  | 414   | _           |
| MJ0765       | 784011  | 785549  | [6Fe-6S] prismane-containing protein {Desulfovibrio desulfiricans}                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             | 603    | 3 5  | 200   | _           |
| Hypothetical |         |         |                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                | 200    | 77.0 | 1339  |             |
| MJ0458       | 1063165 | 1062518 | hypothetical protein (Sulfolobus acidocaldacius)                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               |        |      |       |             |
| MJ0483       | 1047280 | 1048250 | hypothetical protein (Sarcharomyon protein)                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    | -0.9   | -0.9 | 648   | <del></del> |
| MJ0920       | 620866  | 621163  | Control (Section Office) Celevisiae)                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           | 27.7   | 48.7 | 176   | -           |
| M10443       | 1074/80 | /66120  | nypotnetical protein {Mycoplasma genitalium}                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   | 28.3   | 51.3 | 492   | -           |
| Chroma       | 10/4080 | 10/5548 | hypothetical protein {Saccharomyces cerevisiae}                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                | 27.8   | 52.8 | - 699 |             |
| MJ0144       | 1330246 | 1330962 | hypothetical protein (Methanobacterium thermoautotrophicum)                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    | 3,7    | 8    |       |             |
| MJ0044       | 1426552 | 1427241 | hypothetical protein (GP:D38561 6) {Streptomyces wedmorensis}                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  | 5.66   | 0.00 | } {   |             |
| MJ0868       | 680710  | 681000  | hypothetical protein (GP:D63999 31) (Synechocyctis en )                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        | $\top$ | 49.8 | 069   |             |
| MJ1502       | 1662923 | 1663714 | hypothetical protein (GP:D64001 24) (Synechocyetis en 1                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        | 十      | 0.30 | 291   |             |
| MJ1129       | 402152  | 402382  | hypothetical protein (GP:D64001 53) (Sympethocyclis and                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        | $\top$ | 90.1 | 792   |             |
| MJ0057       | 1414899 | 1416176 | hypothetical profein (GP-D64003 25) (C)                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        | 37.5   | 57.9 | 231   |             |
|              |         |         | Special specia | 28.4   | 53.2 | 1278  |             |

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| MJ1335  | 187757  | 187593  | hypothetical protein (GP:D64004 11) (Synechamicia 22)                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          |        |       |      | г             |
|---------|---------|---------|--------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|--------|-------|------|---------------|
| MJ0640  | 902502  | 903458  | hypothetical protein (GP-D64008 43) (c)                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        | 46.2   | 63.5  | 165  | Т             |
| MJ1347  | 177726  | 177280  | hypothetical protein (GD-D44004 24 45                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          | 33.9   | 58.8  | 957  | <del></del> 7 |
| MJ0392  | 1116428 | 1115556 | hypothetical motein (CD-Devote of Synechocystis sp.)                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           | 32.1   | 58.6  | 447  |               |
| MJ0590  | 950234  | 948222  | hypothetical (Or. 104:000 95) {Synechocystis sp.}                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              | 29.1   | 54.3  | 873  |               |
| MJ1178  | 155642  | 366066  | "y Pourciud protein (OP: D04044_18) {Escherichia coli}                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         | 30.6   | 52.6  | 2013 |               |
| MI0438  | 200001  | DCACCC  | hypothetical protein (GP:L47709_14) {Bacillus subtilis}                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        | 27.1   | 55.3  | 315  |               |
| 057057  | 1080099 | 1079128 | hypothetical protein (GP:L47838_15) {Bacillus subtilis}                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        | 29.6   | 8 8 8 | 077  | <del>_</del>  |
| MJ0044  | 898810  | 898223  | hypothetical protein (GP:M18279_1) (Pseudomonas sp.)                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           | 28.3   | 53.4  | 588  |               |
| 0790670 | /23/63  | 723668  | hypothetical protein (GP:M35130_5) {M71467 M71468}                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             |        |       |      | .,.           |
| MJ1526  | 1632280 | 1632810 | hypothetical protein (GP:M36534 1) (Methanohravihagia and 111111)                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              | 78.1   | 2/:1  | 36   |               |
| MJ0888  | 652964  | 653473  | hypothetical protein (GP-1100011 2) (AA                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        | 42.6   | 66.5  | 531  |               |
| MJ0729  | 809665  | 809321  | hypothetical access: (Cr. Cocol 1.2) (Mycobacterium leprae)                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    | 29.5   | 51.4  | 510  | -/,           |
| MJ0787  | 761402  | 750077  | " Pourcite at protein (OP: 018/44_1) {Bacillus firmus}                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         | 29.4   | 56.9  | 345  | 10-           |
|         | 201.07  | //000/  | hypothetical protein (GP:U19363_11) (Methanobacterium thermoautotrophicum)                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     | 49.9   | 71.9  | 1326 |               |
| MJ0693  | 852445  | 853059  | hypothetical protein (GP:1119363 2) (Marth                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     |        |       |      |               |
| MJ0489  | 1039414 | 1038686 | hypothetical profein (GP:1110363 4) (M. A.                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 | 42.8   | 61.9  | 615  |               |
| MJ0446  | 1072662 | 1071784 | honer the first the first that the first the f | 41.3   | 57.5  | 729  |               |
| MJ0076  | 1400741 | 1400403 | "The included protein (GP: 019363_5) {Methanobacterium thermoautotrophicum}                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    | 29.8   | 50.7  | 879  |               |
|         |         | 1400403 | hypothetical protein (GP:U19364_10) (Methanobacterium thermoautotrophicum)                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     | 25.3   | 56.1  | 339  |               |
| MJ0034  | 1435995 | 1436921 | hypothetical protein (GP:U19364 2) {Methanobacterium thermonistics                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             | $\top$ | 7     |      |               |
|         |         |         | $\dashv$                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       | 23.9   | 49.7  | 927  | •             |

| MJ1251   | 277892  | 277392  | hypothetical protein (GP:U19364 4) (Methanohacterium thermoguistical                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          | ;      |          |      |
|----------|---------|---------|-------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|--------|----------|------|
| MJ0927   | 615224  | 615694  | hypothetical protein (GP:1119364 6) /Methanobacterium (Hermitean protein (GP:1119364 6) /Methanobacterium                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     | 37.8   | 91.0     | 201  |
| MJ0785   | 763999  | 762923  | hypothetical protein (GP-1119364 8) (Mathematical protein (GP-111936 8) (Mathematical protein (GP-11936 8) (Mathematical protein | 37.9   | 57.2     | 471  |
| MJ0746   | 799630  | 700036  | (intentational thermosutotrophicum)                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           | 57.5   | 76.6     | 1077 |
| Miller   |         | 55555   | nypotnetical protein (GP:U21086_2) {Methanobacterium thermoautotrophicum}                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     | 60.3   | 76.4     | 306  |
| CCITCIAI | 3/8926  | 380485  | hypothetical protein (GP:U28377_114) {Escherichia coli}                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       | 40.0   | 61.7     | 1560 |
| MJ0653   | 890904  | 890359  | hypothetical protein (GP:U31567_2) {Methanopyrus kandlerit                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    |        |          | 000  |
| MJ0532   | 1003608 | 1004750 | hypothetical protein (GP-1132666 1) that                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      | 7.75   | 8.8      | 546  |
| MJ0674   | 872151  | 871633  | International Control                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         | 39.3   | 59.5     | 1143 |
| 7111500  |         | 671073  | hypothetical protein (GP:X83963_2) {Thermococcus litoralis}                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   | 58.3   | 7.97     | 531  |
| 7551 (M) | 1608984 | 1608592 | hypothetical protein (GP:X85250_3) {Pyrococcus furiosus}                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      | 788    | 009      | 195  |
| MJ0709   | 837195  | 835996  | hypothetical protein (GP:X91006 2) (Pyrococcus sn )                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           |        | 0.00     | 593  |
| MJ0226   | 1255943 | 1255389 | hypothetical profein (GP-740560 1) (Second                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    | 1.67   | 20.5     | 1200 |
| MJ1476   | 25468   | 34051   | (Saccinationity es cerevisiae)                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                | 39.0   | 9.09     | 555  |
|          | 27100   | 16947   | hypothetical protein (HI0380) (Haemophilus influenzae)                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        | 39.7   | 9 63     | 819  |
| MJ0441   | 1076859 | 1076125 | hypothetical protein (H10902) (Haemophilus influenzae)                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        | 十      |          | 2    |
| MJ1372   | 151434  | 150760  | hypothetical protein (H10920) (Haemonhilus in Guine                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           | $\top$ | <u> </u> | 735  |
| MJ0931   | 611416  | 610298  | hybothetical protein (MG372) (M                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               | 46.7   | 67.5     | 675  |
| M10861   | 687740  | 600517  | (injycopiasma genitalium)                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     | 34.9   | 59.9     | 6111 |
|          | 067700  | 088332  | hypothetical protein (MG423) (Mycoplasma genitalium)                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          | 33.9   | 53.9     | 1293 |
| MJ1252   | 77977   | 278609  | hypothetical protein (PIR:B48653) {Lactococcus lactis}                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        | 325    | 133      |      |
| MJ0279   | 1206983 | 1206147 | philies                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       | +      | +        | 653  |
| MJ0299   | 1189620 | 1190600 |                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               | 7.67   | 93.4     | 22   |
|          |         |         | "Transcal protein (TIX.311002) (Thermoplasma acidophilum)                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     | 62.1   | 9.92     | 186  |
|          |         |         |                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               |        |          |      |

| MJ1208                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         | 320842  | 319766  | hypothetical protein (PIR:S21569) (Methanoharterium themselve in the second in the sec |      |      |      | _     |
|--------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|---------|---------|--------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|------|------|------|-------|
| MJ1533                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         | 1625982 | 1627727 | hypothetical protein (PIR-S28724) (Machanical Included Control of  | 55.4 | 74.8 | 1077 |       |
| MJ0323                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         | 1172727 | 1172257 | hypothetical protein (PIR-S38467) (Dec. 16                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     | 67.3 | 83.3 | 1746 |       |
| MJ1162                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         | 368773  | 369060  | hypothetical protein (DID-CALCO), the call protein (DID-CALCO), th | 60.7 | 71.7 | 471  |       |
| MJ0922                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         | 619284  | 619598  | hypothetical action (110,241361) (Methanothermus fervidus)                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     | 48.3 | 67.9 | 288  |       |
| MJ0867                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         | 681124  | 682371  | hypothetical protein (FIR:341383) (Methanothermus fervidus)                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    | 48.6 | 73.4 | 315  |       |
| MJ0047                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         | 1423924 | 1424988 | hypothetical contract (PIR. 2493 / ) (Pseudomonas aeruginosa)                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  | 28.7 | 55.2 | 1248 |       |
| MJ1236                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         | 290570  | 202111  | hand in the state of the state  | 26.9 | 49.9 | 1065 |       |
| MIDIKS                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         | 1306302 |         | hypometical protein (PIR:SS1413) (Saccharomyces cerevisiae)                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    | 33.9 | 54.6 | 1542 |       |
| 7010010                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        | 1300/87 | 1305562 | hypothetical protein (PIR:SS1413) (Saccharomyces cerevisiae)                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   | 32.4 | 56.4 | 1221 |       |
| 8760cM                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         | 614493  | 614957  | hypothetical protein (PIR:S51868) {Saccharomyces cerevisiae}                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   | 787  | 613  |      |       |
| MJ1625                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         | 1535098 | 1533113 | hypothetical protein (PIR:S52522) (Saccharomyces cerentricies)                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 | 70.4 | )    | 465  |       |
| MJ0862                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         | 686185  | 687054  | hypothetical protein (PIR-S52070) (Emilian L.                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              | 27.6 | 50.4 | 1986 | - //. |
| MJ1432                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         | 69872   | 69453   | hymothetical protein (DID Center)                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              | 35.5 | 59.2 | 870  | 2-    |
| MJ0710                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         | 835912  | 834914  | hynothetical protein (CD. Doctoox or control of CD. Doctoox or control | 38.5 | 0.99 | 420  |       |
| MJ0170                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         | 1299322 | 1300185 | hynothetical protein (SD:D1166) (Tr. 1. 1. 1. 1. 1. 1. 1. 1. 1. 1. 1. 1. 1.                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    | 59.2 | 79.9 | 666  |       |
| MJ1593                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         | 1571988 | 1571740 | Comment of the first of the fir | 30.1 | 54.8 | 864  |       |
| MJ0463                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         | 1060122 | 0.000   | etical protein (SP:P12049) {Bacillus subtilis}                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 | 40.3 | 9.69 | 249  | Γ,    |
| M10464                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         | 1000127 | 1050436 |                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                | 78.5 | 92.2 | 309  | C 170 |
| Selective Selection Select |         | CCPCCOI |                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                | 58.8 | 79.4 | 285  | 37 // |
| MJ0130                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         | 1340892 | 1340105 | hypothetical protein (SP:P14027) (Methanococcus vannielii)                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     | 7 63 | +    |      | 1457  |

|                                      |                                                            |                                        |                                          |                                                       |                                                              |                                                     | Г                                                         | T                                                    | $\neg$                                              | Ť                                                  | 13-                                |                                      | Т                                                   | $\overline{T}$                                                                         |                                                     | _                               |                                                          | Т                                                         | _                                                           |                                                     |
|--------------------------------------|------------------------------------------------------------|----------------------------------------|------------------------------------------|-------------------------------------------------------|--------------------------------------------------------------|-----------------------------------------------------|-----------------------------------------------------------|------------------------------------------------------|-----------------------------------------------------|----------------------------------------------------|------------------------------------|--------------------------------------|-----------------------------------------------------|----------------------------------------------------------------------------------------|-----------------------------------------------------|---------------------------------|----------------------------------------------------------|-----------------------------------------------------------|-------------------------------------------------------------|-----------------------------------------------------|
|                                      | 549                                                        | 759                                    | 1263                                     | 1050                                                  | 444                                                          | 423                                                 | 582                                                       | ];                                                   | 2113                                                | 1218                                               | 627                                | 1128                                 | 1434                                                | 3585                                                                                   |                                                     | 399                             | 240                                                      | 267                                                       |                                                             | 506                                                 |
|                                      | 66.3                                                       | 53.9                                   | 67.4                                     | 60.2                                                  | 69.3                                                         | 59.3                                                | 75.4                                                      | 2                                                    | 7.70                                                |                                                    | 54.3                               | 63.3                                 | 52.1                                                | 51.5                                                                                   |                                                     | 62.8                            | 0.09                                                     | 63.9                                                      | 9                                                           | 8.55                                                |
|                                      | 46.9                                                       | 24.1                                   | 45.9                                     | 41.4                                                  | 47.3                                                         | 33.9                                                | 55.9                                                      | 20,2                                                 | 77.66                                               | 42.0                                               | 25.8                               | 38.0                                 | 32.4                                                | 26.9                                                                                   |                                                     | 38.2                            | 40.0                                                     | 42.2                                                      | 30 %                                                        | 20.3                                                |
| hymothetical protein (CD. Diceox, co | hypothetical protein (SD:D16000) (Tethanococcus vannielii) | hypothetical protein (SP-P22340) (Math | hypothetical protein (SD-D2) (1715) (TT- | hypothetical protein (CD: Doctor) (Thermus aquaticus) | hypothetical protein (SF:F23/08) {Methanobacterium ivanovii} | "ypourciteal protein (SP:P28910) {Escherichia coli} | nypotnetical protein (SP:P29202) {Haloarcula marismortui} | hypothetical protein (SP:P31065) (Escherichia coli,) | hypothetical protein (SP:P31466) {Escherichia coli} | hypothetical protein (SP-P11471) (Escharichic 211) | hypothetical acotain (Ch. na. 144) | (St. r.) (St. r.) (Escherichia coli) | hypothetical protein (SP.P31806) {Escherichia coli} | hypothetical protein (SP.P32639) (Saccharomyces cerevisiae) (Saccharomyces cerevisiae) | hypothetical protein (SP:P32698) (Escherichia 2011) | hymothetical action (CD parage) | "yponicular protein (3P:P33382) {Listeria monocytogenes} | hypothetical protein (SP:P33382) {Listeria monocytogenes} | hypothetical protein (SP:P34222) {Saccharomyces cerevisiae} | hypothetical protein (SP:P37002) {Escherichia coli} |
| 1119244                              | +                                                          | 397509                                 | 254676                                   | 75812                                                 | 086360                                                       | \$58070                                             | 9/00/0                                                    | 552658                                               | 1168809                                             | 736679                                             | 1399694                            | 1636646                              | 12/0043                                             | 406336                                                                                 | 450726                                              | 97629                           | 36190                                                    | 070100                                                    | 1419670                                                     | 1635945                                             |
| 1118696                              | 305183                                                     | 398771                                 | 255725                                   | 76255                                                 | 986782                                                       | \$57407                                             | 7.1.63                                                    | 222446                                               | 1170026                                             | 736053                                             | 1398567                            | 1578078                              | 0/00/0                                              | 409920                                                                                 | 451124                                              | 97390                           | 360036                                                   | 202000                                                    | 1419978                                                     | 1636316                                             |
| MJ0388                               | MJ1225                                                     | MJ1133                                 | MJ1273                                   | MJ1426                                                | MJ0549                                                       | MJ0982                                              | MInoon                                                    | 066000                                               | MJ0326                                              | MJ0812                                             | MJ0079                             | MJ1586                               | Y CHILD                                             | 471164                                                                                 | MJ1081                                              | MJ1413                          | M11170                                                   | 1,0001                                                    | I COOCINI                                                   | MJ1523                                              |

| MJ0608 | 934974  | 935750  | hypothetical protein (SP:P37487) {Bacillus subtilie}  | ,;;    |       |      | Г            |
|--------|---------|---------|-------------------------------------------------------|--------|-------|------|--------------|
| MJ1661 | 1493414 | 1493809 | hypothetical protein (SP:P37528) (Bacillus subtilie)  | 44.3   | 71.4  | 111  |              |
| MJ1582 | 1580646 | 1579909 | hypothetical protein (SP:P37545) (Racillus substituti | 47.0   | 72.6  | 396  | <del>-</del> |
| MJ1375 | 148221  | 149408  | hypothetical protein (SP-P17555) (Pacillus auctilis)  | 35.4   | 9.09  | 738  |              |
| MJ0231 | 1249786 | 1250814 | hypothetical protein (SP-P17860) (Pacillus auctilis)  | 25.0   | 48.6  | 1188 | T            |
| MJ0882 | 664582  | 663910  | hypothetical protein (SP-P17873) (Pacillus auxilia)   | 40.0   | 44.0  | 1029 |              |
| MJ0043 | 1429606 | 1427252 | hypothetical protein (SD-D20423) (D                   | 44.0   | 68.7  | 673  |              |
| MJ0048 | 1422159 | 1422842 | hypothetical profession (SP-D28410) (Subtility)       | 45.5   | 58.4  | 2355 |              |
| MJ0989 | 552670  | 553011  | hypothetical protein (SP: P19164) (Escherishing)      | 36.6   | 59.1  | 684  |              |
| MJ1115 | 415733  | 416479  | hymothetical action (CD, Date)                        | 29.0   | \$1.8 | 342  |              |
| M11640 | 1505333 |         | 'y pourciteat protein (SP:P39304) {Escherichia coli}  | 27.1   | 48.3  | 747  | -/           |
|        | 170001  | 120/068 | hypothetical protein (SP:P39587) {Bacillus subtilis}  | 28.9   | 48.5  | 792  | 14.          |
| MJ0577 | 959388  | 958903  | hypothetical protein (SP:P42297) (Bacillus subtilis)  | 7      |       |      | -            |
| MJ0531 | 1004977 | 1004759 | hypothetical protein (SP:P42297) (Bacillus subtilie)  | 0.10   | 20.4  | 486  |              |
| MJ1247 | 282030  | 281677  | hypothetical protein (SP:P42404) (Bacillus embilica   | $\top$ | 68.7  | 219  |              |
| MJ0486 | 1041905 | 1042681 | hypothetical protein (SP-P45476) (Exchanicking        | 7      | 0.09  | 354  |              |
| MJ0449 | 1070080 | 1069565 | hypothetical protein (SD-DA6240) (D                   | 30.6   | 55.7  | 777  |              |
| MJ0682 | 861537  | 864374  | estical process (SE of YOUNG) (Bacillus subtilis)     | 31.8   | 60.7  | 516  |              |
| MJ1677 | 367374  | 200301  | circal protein (Sr.r40830) {Escherichia coli}         | 33.4   | 53.9  | 2838 |              |
|        | 07/0/41 | 0/60/41 |                                                       | 40.3   | 62.0  | 351  |              |
| MJUSSS | 951068  | 952243  | hypothetical protein GP:L07942_2 {Escherichia coli}   |        | 3     |      |              |

| MJ0225   | 1256840 | 1256121 | hypothetical protein GP-1100014 22 (Managed and and and and and and and and and an                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             |                   |            |      |
|----------|---------|---------|--------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|-------------------|------------|------|
| MIDIN    | 1343042 | _       | the state of the s | 27.4 49.0 720     | 49.0       | 720  |
|          | 1342043 | 1342/92 | hypothetical protein GP:U00017 21 (Mycobacterium Jense)                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        | 12.0              |            |      |
| MJ0376   | 1130650 | 0110011 | -pracy                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         | 32.2              | 52.7   750 | 750  |
|          | OCOOC!! |         | hypothetical protein GP:U29579 58 {Escherichia coli}                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           | . 00              |            |      |
| MJ0028   | 1442022 |         |                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                | 30.1 51.5 1521    | 51.5       | 1521 |
|          | 6706441 | 1443844 | hypothetical protein H11305 (Haemophilus influenzae)                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           |                   |            |      |
| Minae    |         |         |                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                | 0./2              | 50.0   822 | 822  |
| 06116141 | 393844  | 394486  | hypothetical protein Lno22n (GP-1141281 22) (C. 1822)                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          |                   |            |      |
| 1110053  |         |         |                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                | 46.2              | 63.8       | 1359 |
| 7CADEMI  | 588063  | 588479  | hypothetical protein PIR:S40613 (Carchagania                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   |                   |            |      |
| 1110403  |         |         | (Saccitationity of Cerevisiae)                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 | 26.8   55.0   417 | 55.0       | 417  |
| [N130403 | 1109067 | 1108276 | hynothetical protein DID Section .                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             |                   |            |      |
|          |         |         | "Jennich Protein FIN. 355199 (Saccharomyces cerevisiae)                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        | 27.6              | 707        |      |
| MJ1031   | 509420  | SORSOR  |                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                | 2,73              | 76/ 7:05   | 76/  |
|          |         | 22000   | Inypometical protein SP:P45869 (Bacillus subtilis)                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             |                   |            |      |
|          |         |         |                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                | × < <             | -          | ~    |

Table 2B

|        | -                                       |
|--------|-----------------------------------------|
|        | adenylate kinase (Methanococons issues: |
|        | 1,049,948                               |
| 000    | 800,000,1                               |
| M10470 | 6/+061vi                                |
|        |                                         |

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Table 3

| <del></del> |       |        |
|-------------|-------|--------|
| MJ0002      | 4071  | 3343   |
| MJ0003      | 4911  | 5378   |
| MJ0008      | 10075 | 10734  |
| MJ0009      | 10743 | 11570  |
| MJ0011      | 12983 | 13459  |
| MJ0012      | 13927 | 1.3427 |
| MJ0013      | 14836 | 14351  |
| MJ0014      | 15455 | 14820  |
| MJ0015      | 15514 | 15804  |
| MJ0016      | 16416 | 15866  |
| MJ0018      | 17658 | 19229  |
| MJ0019      | 21121 | 19232  |
| MJ0021      | 22762 | 23886  |
| MJ0023      | 25284 | 25637  |
| MJ0024      | 26105 | 25689  |
| МJ0025      | 27122 | 26109  |
| МJ0027      | 28572 | 28021  |
| МJ0037      | 38073 | 38786  |
| MJ0038      | 39443 | 38793  |
| MJ0039      | 39974 | 39654  |
| MJ0041      | 41838 | 40477  |
| MJ0042      | 42527 | 41883  |
| MJ0045      | 46506 | 45907  |
| MJ0046      | 47351 | 46569  |
| MJ0050      | 52237 | 51050  |
| MJ0052      | 53374 | 52709  |
| MJ0053      | 54068 | 53388  |
| MJ0054      | 55001 | 54159  |
|             |       |        |

| MJ0056 | 60154  |        |
|--------|--------|--------|
| MJ0062 | 56154  | 55759  |
|        | 60618  | 61238  |
| MJ0063 | 61322  | 61855  |
| MJ0064 | 61897  | 62454  |
| MJ0065 | 63551  | 62463  |
| MJ0066 | 65078  | 63657  |
| MJ0067 | 65160  | 65468  |
| MJ0068 | 65861  | 65517  |
| МJ0070 | 66966  | 67211  |
| MJ0071 | 67211  | 67480  |
| MJ0072 | 67562  | 67693  |
| MJ0073 | 67729  | 68007  |
| MJ0074 | 69089  | 68016  |
| MJ0075 | 70324  | 69236  |
| MJ0077 | 71539  | 70394  |
| MJ0078 | 72674  | 72054  |
| MJ0080 | 74182  | 73802  |
| MJ0086 | 80788  | 81903  |
| MJ0088 | 83019  | 83537  |
| MJ0093 | 88517  | 88092  |
| MJ0094 | 89481  | 88564  |
| MJ0095 | 89828  | 89568  |
| MJ0096 | 90752  | 89967  |
| MJ0100 | 94823  | 93297  |
| MJ0103 | 97958  | 99256  |
| MJ0105 | 101649 |        |
| MJ0106 | 102541 | 101239 |
| MJ0107 | 102733 | 101840 |
| MJ0109 | 106419 | 104295 |
| MJ0110 | 106880 | 105664 |
|        | 100000 | 106614 |

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| MJ0114 | 111874 | 112782 |
|--------|--------|--------|
| MJ0115 | 113249 | 112785 |
| MJ0116 | 113931 | 113257 |
| MJ0119 | 116397 | 115726 |
| MJ0120 | 117070 | 116372 |
| MJ0123 | 119524 | 119195 |
| MJ0125 | 123378 | 123031 |
| MJ0126 | 123685 | 123392 |
| MJ0127 | 124034 | 123672 |
| MJ0128 | 124341 | 124048 |
| MJ0129 | 124487 | 124996 |
| MJ0131 | 126783 | 126475 |
| MJ0133 | 129427 | 128609 |
| MJ0137 | 134976 | 134119 |
| MJ0138 | 136566 | 135121 |
| MJ0139 | 136616 | 138244 |
| MJ0140 | 139150 | 139539 |
| MJ0141 | 139529 | 139825 |
| MJ0142 | 139797 | 140237 |
| MJ0145 | 142991 | 142188 |
| MJ0146 | 143409 | 143203 |
| MJ0147 | 144813 | 143701 |
| MJ0149 | 146003 | 145830 |
| MJ0150 | 146069 | 146587 |
| MJ0154 | 152143 | 152589 |
| MJ0157 | 159807 | 160085 |
| MJ0158 | 160155 | 161276 |
| MJ0159 | 163046 | 161430 |
| MJ0163 | 167378 | 166818 |
| MJ0164 | 168614 | 167430 |
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| MJ0165 | 169394 | 168627 |
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| MJ0173 | 175871 | 176341 |
| MJ0175 | 178089 | 177475 |
| MJ0181 | 182625 | 181918 |
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| MJ0201 | 193486 | 193007 |
| MJ0202 | 193687 | 194454 |
| MJ0206 | 198871 | 198467 |
| MJ0207 | 198967 | 199419 |
| MJ0208 | 200166 | 199429 |
| MJ0209 | 200956 | 200159 |
| MJ0212 | 203759 | 204019 |
| MJ0213 | 204137 | 204583 |
| MJ0215 | 205636 | 205190 |
| MJ0223 | 214474 | 214163 |
| MJ0224 | 215072 | 214566 |
| MJ0227 | 218176 | 219099 |
| MJ0229 | 221136 | 220852 |
| MJ0230 | 221386 | 221144 |
| MJ0233 | 224281 | 225111 |
| MJ0235 | 226124 | 226369 |
| MJ0236 | 226362 | 227639 |
| MJ0239 | 230506 | 230988 |
|        |        |        |

| MJ0240 | 231618 | 231094 |
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| MJ0241 | 232062 | 231628 |
| MJ0243 | 232563 | 232318 |
| MJ0248 | 235142 | 235651 |
| MJ0251 | 238728 | 238288 |
| MJ0252 | 238849 | 239487 |
| MJ0255 | 241359 | 240607 |
| MJ0257 | 242764 | 243696 |
| MJ0258 | 245039 | 243840 |
| MJ0259 | 245717 | 245112 |
| MJ0261 | 247082 | 246423 |
| MJ0263 | 251686 | 250727 |
| MJ0270 | 256421 | 256188 |
| MJ0271 | 256902 | 257441 |
| MJ0272 | 257452 | 257649 |
| MJ0273 | 258107 | 258412 |
| MJ0274 | 260378 | 258819 |
| MJ0275 | 261121 | 260516 |
| MJ0280 | 266375 | 266758 |
| MJ0281 | 267291 | 266761 |
| MJ0282 | 267341 | 267787 |
| MJ0284 | 269902 | 269174 |
| MJ0286 | 270849 | 270499 |
| MJ0287 | 271160 | 270870 |
| MJ0288 | 271755 | 271222 |
| MJ0289 | 272805 | 271801 |
| MJ0290 | 273753 | 273121 |
| MJ0292 | 275409 | 275137 |
| MJ0296 | 279767 | 280360 |
| MJ0297 | 281155 | 280406 |

| MJ0298 | 281290 | 281739 |
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| MJ0301 | 285101 | 284220 |
| MJ0303 | 285971 | 285558 |
| MJ0305 | 286594 | 287778 |
| МJ0306 | 287997 | 287818 |
| MJ0308 | 289084 | 288386 |
| MJ0310 | 290609 | 290268 |
| MJ0311 | 290981 | 290652 |
| MJ0312 | 291845 | 291228 |
| MJ0314 | 293767 | 294369 |
| MJ0315 | 294826 | 294455 |
| MJ0316 | 295458 | 294964 |
| MJ0317 | 296374 | 295733 |
| MJ0319 | 297675 | 297902 |
| MJ0320 | 298001 | 298645 |
| MJ0321 | 298675 | 299040 |
| MJ0325 | 302095 | 301172 |
| MJ0327 | 303625 | 303927 |
| MJ0328 | 304755 | 304318 |
| MJ0329 | 306607 | 304760 |
| MJ0330 | 308266 | 306620 |
| MJ0331 | 308670 | 308266 |
| MJ0332 | 308995 | 308678 |
| MJ0333 | 309670 | 309410 |
| MJ0334 | 309816 | 310112 |
| MJ0335 | 310179 | 310919 |
| MJ0336 | 310932 | 311288 |
| MJ0337 | 311299 | 312084 |
| MJ0338 | 312100 | 312402 |
| MJ0339 | 312374 | 312694 |
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| MJ0340 | 312697 | 313398 |
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| MJ0342 | 313918 | 314286 |
| MJ0343 | 314270 | 316807 |
| MJ0344 | 316820 | 317359 |
| MJ0345 | 317314 | 318264 |
| MJ0346 | 318277 | 318579 |
| MJ0347 | 318593 | 319045 |
| MJ0348 | 319620 | 321995 |
| MJ0349 | 322367 | 322053 |
| MJ0350 | 322681 | 322418 |
| MJ0351 | 323154 | 322705 |
| MJ0352 | 323901 | 323185 |
| MJ0353 | 324142 | 323891 |
| MJ0354 | 324296 | 324123 |
| MJ0355 | 324661 | 324374 |
| MJ0356 | 324957 | 324697 |
| MJ0357 | 326407 | 325943 |
| MJ0358 | 326796 | 326413 |
| MJ0359 | 327449 | 326808 |
| MJ0360 | 328174 | 327770 |
| MJ0361 | 329502 | 329182 |
| MJ0362 | 329659 | 329847 |
| MJ0364 | 332163 | 332495 |
| MJ0365 | 332503 | 333030 |
| MJ0366 | 333033 | 333308 |
| MJ0368 | 334581 | 334886 |
| MJ0369 | 336040 | 334934 |
| MJ0371 | 337418 | 337639 |
| MJ0374 | 339873 | 338884 |
|        |        |        |

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| MJ0377 | 343243 | 343752      |
| MJ0378 | 343921 | 344886      |
| MJ0379 | 345500 | 344889      |
| MJ0380 | 345657 | 345974      |
| MJ0381 | 345977 | 346936      |
| MJ0382 | 346955 | 347683      |
| MJ0383 | 347677 | 349518      |
| MJ0384 | 349546 | 350259      |
| MJ0385 | 350252 | 351304      |
| MJ0386 | 351648 | 351307      |
| MJ0390 | 355149 | 354760      |
| MJ0395 | 357787 | 357314      |
| MJ0398 | 359111 | 359923      |
| MJ0400 | 361593 | 362411      |
| MJ0401 | 362717 | 362520      |
| MJ0402 | 363046 | 362729      |
| MJ0404 | 364804 | 364355      |
| MJ0405 | 365385 | 365002      |
| MJ0408 | 367518 | 367880      |
| MJ0409 | 367946 | 370054      |
| MJ0410 | 370074 | 370865      |
| MJ0414 | 374603 | 373419      |
| MJ0415 | 374712 | 375197      |
| MJ0416 | 375222 | 375791      |
| MJ0417 | 376510 | 375800      |
| MJ0418 | 376627 | 377388      |
| MJ0419 | 377369 | 378430      |
| MJ0420 | 378394 | 379533      |
| MJ0421 | 379640 | 380719      |
|        |        | <del></del> |

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| MJ0423 | 381855 | 382031 |
|--------|--------|--------|
| MJ0424 | 382046 | 382336 |
| MJ0425 | 382317 | 382712 |
| MJ0426 | 383243 | 382704 |
| MJ0427 | 383719 | 383243 |
| MJ0431 | 387350 | 387135 |
| MJ0432 | 388127 | 387852 |
| MJ0433 | 388663 | 388139 |
| MJ0434 | 389342 | 388677 |
| MJ0435 | 389620 | 389342 |
| MJ0437 | 391903 | 391667 |
| MJ0439 | 394280 | 393234 |
| MJ0440 | 394492 | 395292 |
| MJ0444 | 398609 | 397740 |
| MJ0447 | 401037 | 400555 |
| MJ0448 | 401168 | 401935 |
| MJ0450 | 403277 | 403834 |
| MJ0452 | 404962 | 404519 |
| MJ0453 | 405287 | 404967 |
| MJ0455 | 406863 | 406285 |
| MJ0456 | 406888 | 407943 |
| MJ0459 | 410088 | 410354 |
| MJ0480 | 422470 | 423063 |
| MJ0481 | 423792 | 424085 |
| MJ0482 | 423793 | 423074 |
| MJ0485 | 427056 | 428102 |
| MJ0488 | 432390 | 432854 |
| MJ0491 | 434681 | 435106 |
| MJ0492 | 435385 | 435101 |
| MJ0494 | 436499 | 436891 |
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| MJ0496 | 438482   | 438823 |
|--------|----------|--------|
| MJ0497 | 439219   | 438821 |
| MJ0498 | 439679   | 439212 |
| MJ0500 | 442304   | 441537 |
| MJ0501 | 442990   | 442394 |
| MJ0504 | 445785   | 446372 |
| MJ0505 | 446365   | 447117 |
| MJ0512 | 453993   | 453292 |
| MJ0513 | 454868   | 454149 |
| MJ0517 | 459731   | 459321 |
| MJ0518 | 460018   | 459737 |
| MJ0519 | 460275   | 460033 |
| MJ0521 | 461746   | 461549 |
| MJ0522 | 462422   | 461769 |
| MJ0523 | 463226   | 462534 |
| MJ0524 | 463697   | 463239 |
| MJ0525 | 463997   | 463839 |
| MJ0526 | 464308   | 464123 |
| MJ0527 | 465146   | 464655 |
| MJ0528 | 465442   | 465149 |
| MJ0529 | 466215   | 465520 |
| MJ0538 | 474805   | 474026 |
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| MJ1442 | 1412130 | 1412735 |
| MJ1443 | 1412784 | 1413104 |
| MJ1445 | 1414331 | 1414858 |
| MJ1447 | 1415840 | 1416982 |
| MJ1448 | 1416982 | 1418571 |
| MJ1449 | 1418577 | 1419686 |
| MJ1450 | 1419699 | 1420811 |
| MJ1451 | 1420869 | 1422320 |
| MJ1452 | 1422616 | 1423392 |
| MJ1453 | 1423398 | 1423973 |
| MJ1455 | 1425643 | 1424729 |
| MJ1457 | 1427021 | 1427422 |
| MJ1458 | 1427487 | 1428140 |
| MJ1460 | 1430419 | 1429943 |
| MJ1461 | 1431156 | 1430560 |
| MJ1462 | 1431506 | 1431258 |
| MJ1463 | 1432201 | 1431530 |
| MJ1466 | 1436397 | 1435756 |
| MJ1467 | 1436562 | 1437008 |
| MJ1468 | 1437029 | 1440055 |
| MJ1469 | 1440055 | 1440279 |
| МЈ1470 | 1440747 | 1442618 |
| | | |

| | | |
|--------|---------|-------------|
| MJ1471 | 1442618 | 1443151 |
| МЈ1472 | 1443165 | 1444796 |
| MJ1475 | 1446447 | 1446821 |
| MJ1477 | 1447530 | 1448537 |
| MJ1478 | 1449448 | 1448540 |
| MJ1480 | 1451452 | 1452720 |
| MJ1481 | 1452735 | 1453373 |
| MJ1483 | 1454337 | 1454783 |
| MJ1484 | 1454768 | 1455217 |
| MJ1487 | 1459016 | 1460293 |
| MJ1488 | 1460315 | 1461493 |
| MJ1491 | 1465684 | 1466055 |
| MJ1492 | 1466067 | 1466534 |
| MJ1493 | 1466552 | 1467235 |
| MJ1495 | 1468532 | 1469377 |
| MJ1496 | 1469370 | 1469711 |
| MJ1497 | 1469711 | 1470748 |
| MJ1499 | 1472128 | 1471649 |
| MJ1500 | 1472920 | 1472363 |
| MJ1501 | 1473615 | 1472947 |
| MJ1503 | 1474982 | 1474587 |
| MJ1506 | 1479963 | 1478767 |
| MJ1507 | 1480030 | 1481214 |
| MJ1509 | 1482024 | 1482482 |
| MJ1510 | 1483084 | 1482506 |
| MJISI1 | 1483234 | 1483572 |
| MJ1513 | 1489601 | 1488606 |
| MJ1514 | 1489692 | 1490078 |
| MJ1515 | 1490084 | 1491148 |
| MJ1516 | 1491173 | 1491466 |
| | | |

| MJ1517 | 1492030 | 1492863 |
|--------|---------|---------|
| MJ1518 | 1492917 | 1493975 |
| MJ1519 | 1494094 | 1497618 |
| MJ1520 | 1498588 | 1497656 |
| MJ1521 | 1498905 | 1500170 |
| MJ1524 | 1501404 | 1501727 |
| MJ1525 | 1501702 | 1504500 |
| MJ1527 | 1505607 | 1505281 |
| MJ1535 | 1512870 | 1513766 |
| MJ1537 | 1515742 | 1514714 |
| MJ1539 | 1516728 | 1517042 |
| MJ1540 | 1517209 | 1517466 |
| MJ1542 | 1521169 | 1518746 |
| MJ1544 | 1523759 | 1522470 |
| MJ1545 | 1523900 | 1524592 |
| MJ1547 | 1525820 | 1526005 |
| MJ1548 | 1526062 | 1526427 |
| MJ1550 | 1527849 | 1528031 |
| MJ1551 | 1528046 | 1528216 |
| MJ1553 | 1528749 | 1529240 |
| MJ1554 | 1529326 | 1531191 |
| MJ1556 | 1532701 | 1533636 |
| MJ1557 | 1533644 | 1534390 |
| MJ1558 | 1534666 | 1534397 |
| MJ1559 | 1534699 | 1535262 |
| MJ1561 | 1538168 | 1536510 |
| MJ1562 | 1539331 | 1538168 |
| MJ1563 | 1539812 | 1539345 |
| MJ1564 | 1540186 | 1540695 |
| MJ1565 | 1540699 | 1542237 |
| | | |

| | MJ1566 | 1543572 | 1542232 |
|---|--------|---------|---------|
| | МЈ1567 | 1544072 | 1543557 |
| | MJ1568 | 1544632 | 1544078 |
| | MJ1570 | 1545637 | 1545981 |
| | MJ1571 | 1546111 | 1546986 |
| | MJ1573 | 1548452 | 1548270 |
| | MJ1576 | 1551559 | 1552164 |
| | MJ1577 | 1552197 | 1553990 |
| | MJ1579 | 1555146 | 1554937 |
| | MJ1580 | 1555498 | 1555127 |
| | MJ1583 | 1557431 | 1557808 |
| | MJ1584 | 1558268 | 1557816 |
| | MJ1585 | 1559172 | 1558255 |
| | MJ1587 | 1560732 | 1561265 |
| | MJ1588 | 1561285 | 1561620 |
| | MJ1589 | 1561657 | 1562379 |
| İ | MJ1590 | 1562770 | 1563084 |
| | MJ1595 | 1567357 | 1566332 |
| | MJ1598 | 1572075 | 1571026 |
| | MJ1599 | 1572924 | 1572094 |
| | MJ1600 | 1573002 | 1573532 |
| | MJ1601 | 1573539 | 1574018 |
| | MJ1604 | 1578693 | 1577308 |
| | MJ1608 | 1582917 | 1583126 |
| | MJ1609 | 1583168 | 1584289 |
| | MJ1613 | 1589822 | 1589058 |
| | MJ1614 | 1590582 | 1589830 |
| | MJ1615 | 1591350 | 1590586 |
| 1 | MJ1617 | 1593103 | 1593381 |
| | MJ1618 | 1593786 | 1593397 |
| | | | 1 |

| MJ1620 | 1594531 | 1596084 |
|--------|---------|---------|
| MJ1621 | 1596297 | 1596127 |
| MJ1622 | 1597169 | 1597719 |
| MJ1623 | 1597939 | 1599474 |
| MJ1624 | 1599991 | 1599602 |
| MJ1626 | 1602381 | 1600087 |
| MJ1627 | 1604683 | 1604231 |
| MJ1628 | 1606127 | 1604784 |
| MJ1629 | 1607293 | 1606418 |
| MJ1630 | 1610737 | 1607330 |
| MJ1631 | 1611184 | 1612740 |
| MJ1632 | 1612697 | 1613446 |
| MJ1633 | 1614897 | 1613467 |
| MJ1634 | 1615733 | 1615011 |
| MJ1635 | 1615933 | 1617174 |
| MJ1637 | 1618268 | 1619686 |
| MJ1638 | 1620457 | 1619678 |
| MJ1639 | 1620605 | 1621036 |
| MJ1640 | 1621671 | 1621057 |
| MJ1641 | 1622664 | 1621804 |
| MJ1642 | 1623032 | 1623514 |
| MJ1644 | 1627146 | 1627667 |
| MJ1646 | 1628442 | 1629074 |
| MJ1650 | 1632586 | 1631435 |
| MJ1651 | 1633407 | 1632631 |
| MJ1653 | 1635797 | 1636951 |
| MJ1654 | 1637097 | 1637693 |
| MJ1657 | 1639687 | 1640427 |
| MJ1658 | 1640511 | 1640783 |
| AJ1659 | 1640800 | 1641870 |

| MJ1660 | 1641857 | 1643503 |
|---------|---------|---------|
| MJ1664 | 1646502 | 1647179 |
| MJ1665 | 1648555 | 1647182 |
| МJ1666 | 1650080 | 1648686 |
| MJ1667 | 1651336 | 1650083 |
| MJ1668 | 1652321 | 1651194 |
| MJ1669 | 1653119 | 1652376 |
| MJ1670 | 1653547 | 1653149 |
| MJ1671 | 1653684 | 1653550 |
| MJ1672 | 1656206 | 1653807 |
| MJ1673 | 1656630 | 1656244 |
| MJ1674 | 1658539 | 1656638 |
| MJ1676 | 1659621 | 1660334 |
| MJ1678 | 1660939 | |
| MJ1679 | 1662142 | 1662126 |
| MJ1680 | 1662411 | 1662432 |
| MJ1681 | 1663887 | 1662866 |
| MJECS01 | 1268 | 1662862 |
| MJECS02 | 4814 | 432 |
| MJECS03 | 5192 | 1272 |
| MJECS04 | 5884 | 4851 |
| MJECS05 | | 5459 |
| MJECS06 | 6365 | 6814 |
| MJECS07 | 7443 | 7009 |
| MJECS08 | 8765 | 7428 |
| MJECS09 | 11950 | 8738 |
| | 12641 | 11925 |
| MJECS10 | 14062 | 13181 |
| MJECS11 | 14404 | 15030 |
| MJECS12 | 16547 | 15411 |
| MJECL01 | 275 | 1048 |

| MJECL02 | 1474 | 1085 |
|---------|-------|-------|
| MJECL03 | 1700 | 1377 |
| MJECL04 | 1865 | 3250 |
| MJECL05 | 3235 | 3450 |
| MJECL06 | 4170 | 3787 |
| MJECL07 | 5844 | 4561 |
| MJECL08 | 7415 | 5832 |
| MJECL09 | 7780 | 8103 |
| MJECL10 | 8107 | 8784 |
| MJECL11 | 8788 | 9159 |
| MJECL12 | 9150 | 9887 |
| MJECL13 | 10678 | 12483 |
| MJECL14 | 14468 | 15427 |
| MJECL15 | 15420 | 16541 |
| MJECL16 | 16599 | 16811 |
| MJECL18 | 20873 | 21505 |
| MJECL19 | 21456 | 22019 |
| MJECL20 | 22829 | 23290 |
| MJECL21 | 24596 | 23298 |
| MJECL22 | 25120 | 24854 |
| MJECL23 | 27628 | 25136 |
| MJECL25 | 28835 | 29167 |
| MJECL26 | 30215 | 29178 |
| MJECL27 | 31077 | 30571 |
| MJECL28 | 35352 | 31534 |
| MJECL30 | 37621 | 37151 |
| MJECL31 | 37811 | 37599 |
| MJECL32 | 40153 | 38828 |
| MJECL33 | 41381 | 40125 |
| MJECL34 | 43121 | 42231 |
| | | |

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| MJECL35 | 45007 | 43115 |
|---------|-------|-------|
| MJECL36 | 45921 | 45394 |
| MJECL37 | 46065 | 46865 |
| MJECL38 | 47997 | 47197 |
| MJECL39 | 49387 | 48329 |
| MJECL41 | 53908 | 52613 |
| MJECL43 | 57371 | 56187 |
| MJECL44 | 58339 | 57341 |

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Table 4

| | Genes of M. jannaschii that contain inteins. | |
|-------------|---|----------------|
| Gene
No. | Putative identification | No. of inteins |
| MJ0043 | Hypothetical protein (Bacillus subtilis) | 1 |
| MJ0262 | Putative translation initiation factor, FUN12/IF-2 family | + |
| MJ0542 | Phosphoenolpyruvate synthase | +-: |
| MJ0682 | Hypothetical protein (Escherichia coli) | + |
| MJ0782 | Tranascription initiation factor IIB | 1 |
| MJ0832 | Anaerobic ribonucleoside-triphosphate reductase | 2 |
| MJ0885 | DNA-dependent DNA polymerase, family B | 2 |
| MJ1042 | DNA-dependent RNA polymerase, subunit A' | 1 |
| MJ1043 | DNA-dependent RNA polymerase, subunit A" | , |
| MJ1054 | UDP-glucose dehydrogenase | 1 |
| MJ1124 | Hypothetical protein (Saccharomyces cerevisiae) | |
| MJ1420 | Glutamine-fructose-6-phosphate transaminase | |
| AJ1422 | Replication factor C, 37-kD subunit | 3 |
| /J1512 | Reverse gyrase | |

PCT1.WPD

The 1,664,976 M. jannaschii circular chromosome (SEQ ID NO:1) has the following sequence:

GGATTATTATGCTACTGGTTTTAAAATAATTGACTTATCTAAACTAAAAGGAGGAATTAA 5 **GGATTTAAAATAAATTCGCTTATCTTCTCTTCAATTTTTATTACTCATAAAAATTA** GTCTGGGAATAAAAACCAAAATTGCCCAAAATGTAATAACAGCCCATGGATACAAAGAGC AAATAATTTTATTGCTCAAAATCAAAATGTTCAAACAGGTACTAAGGAATATTATCAAGT TGAAGCAGTAAAGTACTTATTAAATAATGGACATTGTGGGATAGATTGTAGGGCAAAAAT 10 TAGCGATATTATAAAGGGAATAAATTATCCCAAAAATAGGGAAGCTTTCCAACATGAAGT GTTGATACCACTAAAACAGTATGGCATCATAGCAACATTGGTTTATCCAGGACGTAAAGG AGGCGTATTTATCCCATGTAATAATGATGAAATAAAAAAAGTGGCAAAACAAGTGTTTAA GAGGATAGAAAGTGAATTAGAAAATTTAGAAGGTTCTGCGACAGGAGTTCAAAATATAAA 15 TGCATCAAGAGTAATTATGTTTTTGTTTTTTACATTATCAAATTTTCCATCTGTTTTTAA AAGTTCTTTTTTTTTCCTCTCTCTCCAACTCTGCAATAGTATTCATCAATCTCAAAGCC **AATATAATCAATCCCTAACCTAATACATGCTATTGCTGTGCTTCCAATTCCCATAAATGG** GTCTAAAACAAGATTTGTCTTTTTAACACCATGCAATTTAATACACATCTCCGGAAGTTT TGGAGGAAATGTTGCAGGATGAGGTCTTTCTTTTTCTTTTGATTGGATTGTTTCATAAGG 20 GATAAACCACGTATTTCCCCTATCTCTTAAATCTCCTTTTCTGTTAAATCTCTTTATATT GCTTTTATCCTGATAAGGAACACCAATTGCTAATTTGTCTAACTTAACGTTCCCATTTTT TGTGAAGTGGAAAATATATTCATGCATTATACTTAAAAATCTATCACTGTTTATTGGCTT GTNATGTCCAACAGCAATATCTCCAATAATATTTGGGTAATTTCCAACATCTTCTTTTTG TATTGCAATTGATTTTACCCAATGTATAGTATTTTGTAATTTTAAAATGTTTTCTTATAAC 25 ATTAGCAACATCAAAGGCAATCCACGGGTCTTTTGCAGTATAGCCAACATTTATAAAAAA TGAGCCGTCATCTTTAATACTCTCTTTATTTCTTTGACAACTTCTTCAATCCAATTTAA **ATAATCTTCTCTACTTAAATTATCAGAGTATTTGTTGTTATTTTATGCCAATATTATAGGG** TGGAGACGTAACAACAACAACTGTCTTATCTTTTAACTGTTTCATTCCCTCTAAACA ATCCATACAGTAGATTTTATTTATCTCCATTTTTAATCCCCATCATTATTTATTCTATCA 30 TCAATTCTGCAAGCTTCTCTACTTCTTTAATTCCCCTATCAAAATCATTTAAGTTTAAAT TTAAAAATTTGTTGAAGTAATATGCTTCGCAAGTAGCATTAAAAAATGATATTTTAAAGT GCTTAGACAACTTATTTATTAACTCTTTATTTTCAAGCATGTAGAAATTAGCATAATGTC TTTCAGGATTTAATGAGCTTTTTATATGCTTTGAATAATTTTTTTGAGATAAAAGTCGT 35 CTATCTTTTTTATTATATCTTTTTCAACACTTCTAACATCAAATAAGACATAAGCATAAT CTGGAATGATATTGCTTTGAATTCCTCCTTTTATTATGGTTGGAGTTATTGAAGAACTGT AGATTTTATCAACCTTAATCTTTTCCAAAGGAAGATTTTTTAAATCTAAAATAACTCTGC TTAAGATTTCTATTGGATTTAGGCCTTGAGATGAGGCATGCCTCGCCTCCCCAAAACTTT CAACAATATACTCAAATCTTCCTTTATGTCCAATACAAACATTTAAGTCAGTAGGCTCTC 40 TGTAAATACCATTTGATTCTGTTTCTTCATCAGGAGATATAACTAATAGAGAGTTATTGC TATTTAAAAAGCATGAATCATTAAAACCACATTCCCTTTAGCATCTATAACTCCAGTCC CATAAAAATTGTTATCATCTTTTTTAAAATTTGATTGAATCTTTACAGTGTCTATATGTG **AATTTAATATCAAATCAAAGTTTTCTTTTTTTTTATATGCTACAAAGCATCCTTCAATGA** 45 TAGTATTTTTTATTCCTAAGTTATTGAAAAGATTAGATAAATATTTAAATGCCTTTTTAA CACCAATTCTATTATCCGTCCTAATTTTCACCAAATCCTCTAAGATTTTTAAATAATCCA TAATTATCATCTCATAAATTCTACTTTTTCTCCAATAATTTCATTTAAATCAATATCACT ACACTTAAATTCAAGCATTGCTGTTGAGTAATTTTTACATTTGTAGGTTTTCCATGGCTT TAATCTTACAGCTTCGACAACCCTATTTTTATCAATAAAATTATATCAATAGGATAAAG 50 TCCAATATCTCTAAGCATTAAACCAAAAGCTCTTTTAATAAAATTATCTGCCAATACAAC TTCAAATTCTAAATTTCCAACTTTAACTTTTTTAATTTTCTTATTTTGCATTTTTTCAC TTTCTTTTTTGCTGTATGGGACAGGGATGTAATAAACTGAAGGTTTGGCTCCCATTGGTT GTGGATAAAGCTCTAATAACTCATAAACCTTTCTTGGAACATTTGTATTAACTTCAATAC 55 CTAATTCTTTTAATTTACTAACTGTTAAAGGGTAATCATGTGTCCATGTTCCTGAAGTTA GTTTTTTTGCGATTTCTTTAGCTTTTTCATCTCCATATTTATCTTTCAACAACTCATAAA CAAATTCTTCCATCTGTTTAATAGCTTTTTTAGATATATCAACCAATATTAATGTCTCAT CACTTACTTTTTCTCCCTTCCTATAGTATGCCTCTAAGATaGATGCAGCAGGATACTGCC CAATCTGTGGATCTACTGGCCCCATTACAGCGTTTTTATCCATAATTATTTCATCTGCAG 60 CTAAGGCAATTAAACTTCCTCCACTCATCGCATAATGTGGAATTATAACTGTTGTTTTTG CCTTATGTTCCTTTAAAGCTAAGGCTATCTGCTCACTCGCTAAAGCTAAACCTCCAGGAG TATGAATGATTAAATCAATAGGCATATCTTCTGGTGTTAATCTAATAGCCCTCAAAATCT CTTCACTATCTTCAATAGTGATAAATTTATATATTTGGTATCCCTAAGAATGTTAATGCTT CTTGTCTATGTATCATAGCTATAACTCTTGTTCCCCTCTGTCTTTCAATCTCCCTTATAC 65

| | TTTTTATTTATAAAATTCAAAAAATATCTTATCGTATTATAGAAAGATTTGTAATA |
|-----|--|
| | AACTCATTATAATAGTGAAATCTTACTTCGAAAATTTCTAACCTTGGCTGAACTTTGTTG |
| | ATTAAGTTCAGGATAAACAAAAAATAAAAAGAACAATGATTTTAAACTCACTATCAGTGT |
| _ | AGAGATTGGCATTAACTATTTATTTGTATTTATCTATCATACTGAGAGAGTTTTTTTATTTT |
| 5 | CTTTTATTGCTTTATTGATTTTTCTTTGAATGATTCTAGTACTATTTTCTCATAAGGAAA |
| | ANTGTTTGGTTTGTCATCTTAAATTTAAGTGATTTGATAAATTATAATTATCCCAACTTA |
| | AACTGTAAATGAACTACAATATCCTTTCCTTTTTGTTTAAGTTCTATATCTTTTATTTTT |
| | GAACAATTTCCACAGAATTCTTTTCTTAATATGTTTTTTATGTATCGGCATAAAGATTCT |
| | TTGATTATTGCATCGTTTATATCATACCAAATTGCATAATTTTTGAGTTCGAATTCAAAA |
| 10 | TTTGGCTTTTTACTCTTCATTACCTCATATATTTCTTTAATAATA |
| | TTAATATATTCTTCATTTAATAAGTTTTTATCGATATATTTTTCAATATTTTCTTTC |
| | GAGTCAAAACCATTTTCTTTCGTATTATTTTATAGATTTTATACCCTCTTCCTT |
| | CTATGGTTAGTGGATTGTATTTTTATCCATGTGTTAATTTCACGAATGGGTAGTCGTTGG |
| | TCTATTACATAGATTTTTCCATTTATCTCAATAGCTGCAGCAACATGCATAGGATGAGTT |
| 15 | ACTAAATATAAGTTATAGTTGGGAAACAAATTCGAAAGTAAAGCAATAGTTAGCTTAGCA |
| | TAATCTCTACATACTGCTTTTTTGTATTTTAGAATTTTTGGACACTTTAATATCATAACAT |
| | AAAGTATCATGCAGCATAGCGATTGCTGATACAAAATTTCCATTGCATTGAAACAAATAT |
| | TTTACTATCAATGTAAGTATTAACACCCAAAGTCCTGAAATTATACTAATAATTATAAGA |
| • • | TTTTCACTGAAAATATACAATGTGAAGATACTTACTAACAACAACATGCTAATGTTGTTT |
| 20 | AAGTATTGTGAATATTTTGATGGCAATGATATTAATACTAAGAGAGCCACTACTGCAGAT |
| | ATTACCCATAGTATCAATACCATAATATCATTGATTATAATCTCAAAACCTATTATCAAT |
| | AACAATACCATAAATAACAATACCACACCATATAACATAGCCGCAATAACATAATAAATT |
| | AAAGAATCTGCCGCTCTTTCCATCCAATATCTAATATTAGTTTCTTGCCATTCCAAAATA |
| | TTATTTAAAGTTTCAACAATTGAATTTTCCCATAACTGTTCAGACAGTTTTTTATTTCG |
| 25 | TTACTATAAATTTCTTTTAGAGAAGGAATACTTAAAAATGTGTGACAACTT |

While the present invention has been described in some detail for purposes of clarity and understanding, one skilled in the art will appreciate that various changes in form and detail can be made without departing from the true scope of the invention.

All patents, patent applications and publications recited herein are hereby incorporated by reference.

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What Is Claimed Is:

- 1. An isolated nucleic acid molecule comprising a polynucleotide having a nucleotide sequence at least 95% identical to a sequence selected from the group consisting of:
- (a) a nucleotide sequence of an open reading frame depicted in Table 2(a) or 3;
- (b) a nucleotide sequence of an open reading frame depicted in Table 2(a) or 3, but minus the codon for the N-terminal methionine residue, if present; and
- (c) a nucleotide sequence complementary to any of the nucleotide sequences in (a) or (b).
- 2. An isolated nucleic acid molecule comprising a polynucleotide having a nucleotide sequence 100% identical to a sequence in (a), (b) or (c) of claim 1.
- 3. An isolated nucleic acid molecule comprising a polynucleotide that hybridizes under stringent hybridization conditions to the nucleic acid molecule of claim 2.
- 4. An isolated nucleic acid molecule comprising a polynucleotide that encodes the amino acid sequence of an epitope-bearing portion of the *M. jannaschii* protein encoded by an open reading frame depicted in Table 2(a) or 3.
- 5. A method of making a recombinant vector comprising inserting the isolated nucleic acid molecule of claim 1 into a vector.
 - 6. A recombinant vector produced by the method of claim 5.

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- 7. A method of making a recombinant host cell comprising introducing the recombinant vector of claim 6 into a host cell.
 - A recombinant host cell produced by the method of claim 7.
- 9. A recombinant method for producing a *M. jannaschii* polypeptide, comprising culturing the recombinant host cell of claim 8 under conditions such that said polypeptide is expressed and recovering said polypeptide.
- 10. An isolated polypeptide having an amino acid sequence at least 95% identical to the amino acid sequence selected from the group consisting of:
- (a) an amino acid sequence encoded by a M. jannaschii open reading frame depicted in Table 2(a) or 3; and
- (b) an amino acid sequence encoded by a *M. jannaschii* open reading frame depicted in Table 2(a) or 3, but lacking the N-terminal methionine residue.
- 11. An isolated polypeptide, wherein said amino acid sequence is 100% identical to a sequence in (a) or (b) of claim 10.
- 12. An isolated antibody that binds specifically to the polypeptide of claim 11.
- 13. Computer readable medium having recorded thereon the nucleotide sequence depicted in SEQ ID NO:1, 2, or 3, or a nucleotide sequence at least 99.9% identical thereto.
- 14. Computer readable medium having recorded thereon the nucleotide sequence of at least one *M. jannaschii* open reading frame depicted in Table 2(a) or 3 or its complement.

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- 15. The computer readable medium of claim 13, wherein said medium is selected from the group consisting of a floppy disc, a hard disc, random access memory (RAM), read only memory (ROM), and CD-ROM.
- 16. The computer readable medium of claim 14, wherein said medium is selected from the group consisting of a floppy disc, a hard disc, random access memory (RAM), read only memory (ROM), and CD-ROM.
- 17. A computer-based system for identifying fragments of the *M. jannaschii* genome that are homologous to target nucleotide sequences, comprising:
- (a) a data storage means comprising the nucleotide sequence of SEQ ID NO:1, 2, or 3, or a nucleotide sequence at least 99.9% identical thereto;
- (b) a search means for comparing a target sequence to said nucleotide sequence of said data storage means of step (a) to identify a homologous sequence, and
- (c) a retrieval means for obtaining said homologous sequence of step (b).

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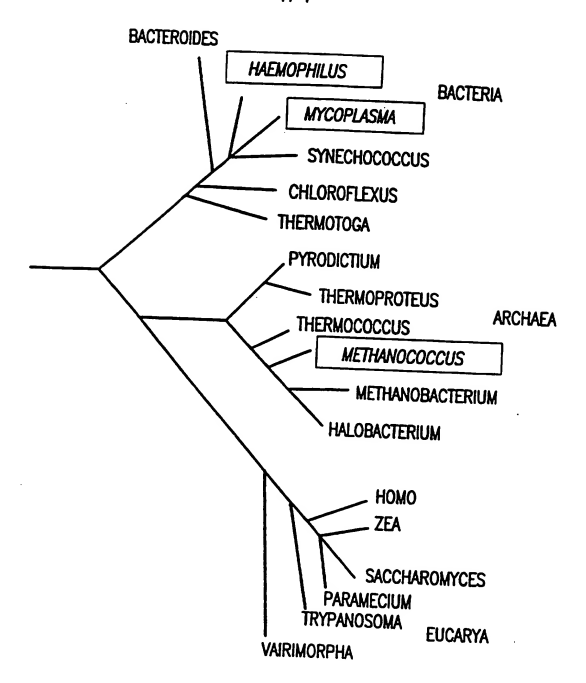
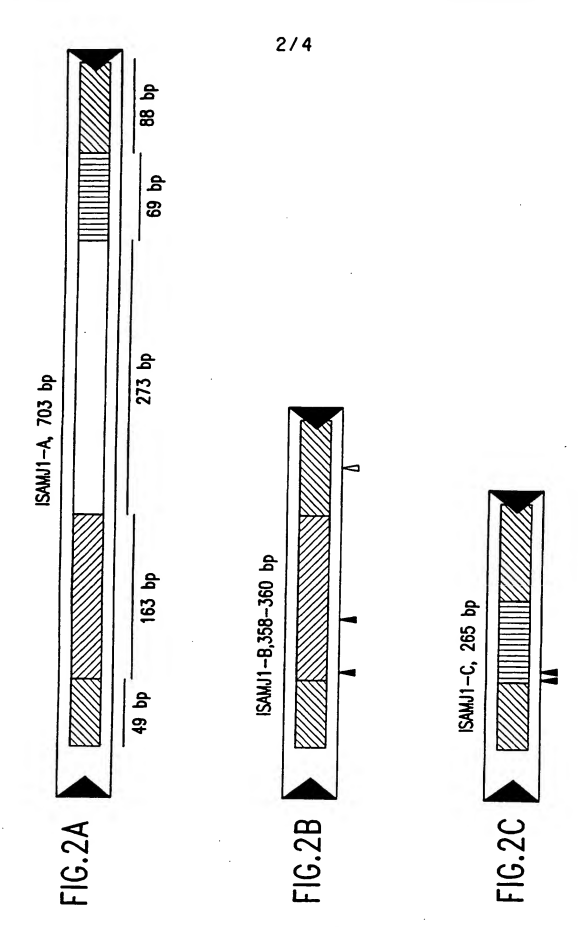
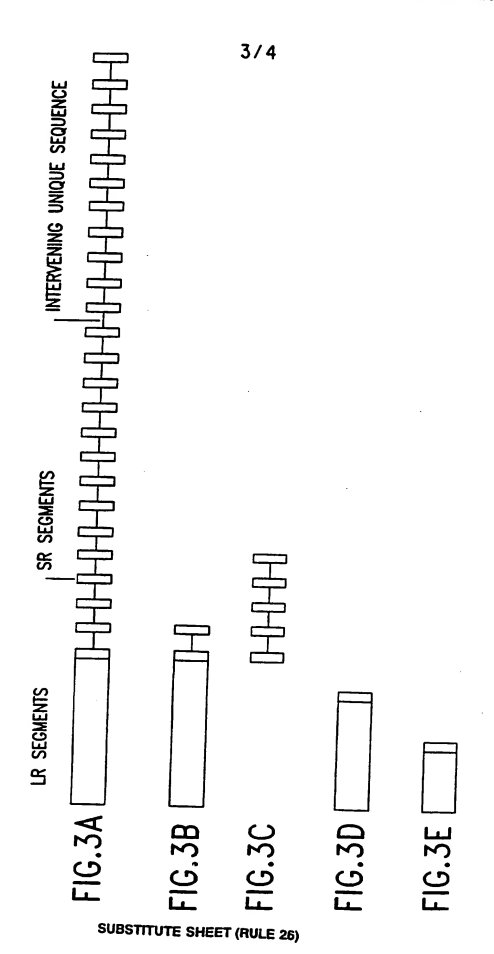


FIG.1

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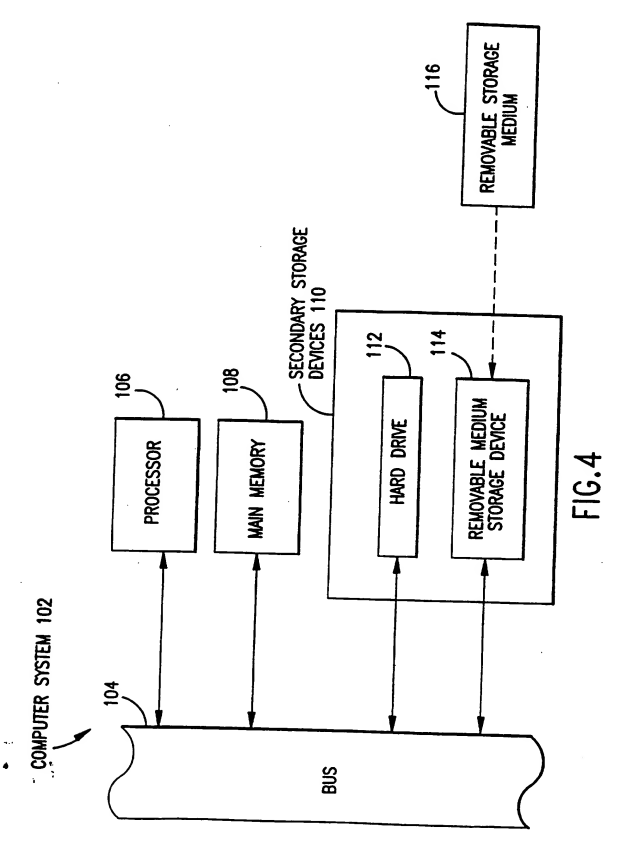


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(88) Date of publication of the international search report: 26 March 1998 (26.03.98)

(54) Title: COMPLETE GENOME SEQUENCE OF THE METHANOGENIC ARCHAEON, METHANOCOCCUS JANNASCHII

(57) Abstract

The present application describes the complete 1.66-megabase pair genome sequence of an autotrophic archaeon, Methanococcus jannaschii, and its 58- and 16-kilobase pair extrachromosomal elements. Also described are 1738 predicted protein-coding genes.

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